

pharmacies involved in the treatment of PAH and PH-ILD in the United States, as well as key stakeholders involved in the distribution and reimbursement of medicines to treat these patients. We established our commercial presence in the field to support Treprostinil Injection and have since expanded our presence to support the potential launch of YUTREPIA (treprostinil) inhalation powder (â€œYUTREPIAâ€), further validating our reputation as a company committed to supporting PAH and PH-ILD patients. We conduct research, development and manufacturing of novel products by applying our subject matter expertise in cardiopulmonary diseases and our proprietary PRINTÂ® technology, a particle engineering platform, to enable precise production of uniform drug particles designed to improve the safety, efficacy and performance of a wide range of therapies. Through development of our own products and research with third parties, we have experience applying PRINT across multiple routes of administration and drug payloads including inhaled therapies, vaccines, biologics, nucleic acids and ophthalmic implants, among others. Our lead product candidate is YUTREPIA for the treatment of PAH and PH-ILD. YUTREPIA is an inhaled dry powder formulation of treprostinil designed with PRINT to improve the therapeutic profile of treprostinil by enhancing deep lung delivery while using a convenient, low effort dry-powder inhaler (â€œDPIâ€) and by achieving higher dose levels than the labeled doses of current inhaled therapies. On August 16, 2024, the United States Food and Drug Administration (the â€œFDAâ€) (i) granted tentative approval for our New Drug Application (â€œNDAâ€) for YUTREPIA for the treatment of PAH and PH-ILD and (ii) simultaneously determined that Tyvaso DPI, approved on May 23, 2022, qualifies for a three-year New Clinical Investigation exclusivity for the chronic use of dry powder formulations of treprostinil for the approved indications. As a result, final approval of YUTREPIA for PAH and PH-ILD is currently delayed until after expiry of the three-year regulatory exclusivity for Tyvaso DPI on May 23, 2025. We are also developing L606, an investigational, liposomal formulation of treprostinil administered twice-daily with a short-duration next-generation nebulizer, which we licensed from Pharmosa Biopharm Inc. (â€œPharmosaâ€). L606 is currently being evaluated in an open-label study in the United States for treatment of PAH and PH-ILD with a planned pivotal study for the treatment of PH-ILD. Risks and Uncertainties We are subject to risks and uncertainties common to companies in the biotechnology industry, including, but not limited to, development by competitors of new technological innovations, dependence on third parties and key personnel, protection of proprietary technology, compliance with government regulations, and the ability to secure additional capital to fund operations.â€¹⁰ Table of Contents The current global macro-economic environment is volatile, which may result in supply chain constraints and elevated rates of inflation. In addition, we operate in a dynamic and highly competitive industry and believe that changes in any of the following areas could have a material adverse effect on our future financial position, results of operations, or cash flows: the ability to obtain future financing; advances and trends in new technologies and industry standards; results of clinical trials; regulatory approval, market acceptance and third-party payor coverage for our products; development of sales channels; certain strategic relationships; litigation or claims against us, including claims related to intellectual property, product, regulatory, or other matters; and our ability to attract and retain employees necessary to support our growth.â€ Product candidates we develop require approval from the FDA and/or other international regulatory agencies prior to commercial sales. There can be no assurance that our product candidates will receive the necessary approvals or, if we do, the indications for which our products will be approved. If we are denied approval, approval is delayed, approval is for less than all of the indications we are seeking, or we are unable to maintain approval, it could have a material adverse impact on our business, financial position and results of operations.â€ We rely on single source manufacturers and suppliers for the supply of our product candidates, adding to the manufacturing risks we face. In the event of any failure by a supplier, we could be left without backup facilities. Any disruption from these manufacturers or suppliers could have a negative impact on our business, financial position and results of operations.â€ Liquidity We have financed our growth and operations through a combination of funds generated from revenues, the issuance of convertible preferred stock and common stock, bank borrowings, bank borrowings with warrants, the issuance of convertible notes and warrants, and revenue interest financing. Since inception, we have incurred recurring losses, including a net loss of \$92.0 million for the nine months ended September 30, 2024. As of September 30, 2024, we had an accumulated deficit of \$521.1 million.â€ We expect to incur significant expenses and operating losses for the foreseeable future as we advance our product candidates through clinical trials, seek regulatory approval of such product candidates and pursue commercialization of any approved product candidates. These efforts require significant amounts of capital, adequate personnel and infrastructure, and extensive compliance-reporting capabilities. Even if our development efforts are successful, it is uncertain when, if ever, we will realize significant revenue from product sales. Additionally, the Revenue Interest Financing Agreement with HealthCare Royalty Partners IV, L.P. (â€œHCRâ€) dated January 9, 2023, as amended (the â€œRIFAâ€) contains fixed quarterly payments and minimum cash covenants that require us to maintain cash and cash equivalents in an amount at least equal to \$7.5 million during the calendar year beginning on January 1, 2024 and at least equal to \$15.0 million for the remainder of the payment term after the calendar year ended December 31, 2024.â€ In September 2024, we entered into the Fifth Amendment to the RIFA pursuant to which HCR funded an additional \$32.5 million on September 12, 2024, for total funding of \$100 million. Additionally, payments due under the RIFA were amended such that the one-time fixed payment previously due in July 2025 is now due in equal payments in January and July 2026. See Note 12A for further information.â€ On September 12, 2024, we sold shares of our common stock in an underwritten registered public offering for net proceeds of approximately \$53.7 million and sold shares of our common stock for net proceeds of approximately \$10.0 million in a private offering. See Note 8 for further information.â€ Our future funding requirements will be heavily determined by the timing of the potential commercialization of YUTREPIA and the resources needed to support development of our product candidates. Based on our current plans, we expect that we will require additional capital to fund operations as well as to pursue in-licenses or acquisitions of other product candidates. If we are unable to obtain additional funding, we could be required to delay, reduce, or eliminate research and development programs, product portfolio expansion, or future commercialization efforts, which could adversely affect our business prospects, or we may be unable to continue operations.â€¹¹ Table of Contents Although we expect to continue to generate operating losses for the foreseeable future, we believe that as a result of the recent Fifth Amendment to the RIFA and net proceeds from the sale of common stock described above, excluding any future YUTREPIA product revenue, our cash and cash equivalents will be sufficient to fund operations, capital expenditures, and RIFA payments and allow us to remain in compliance with our minimum cash covenants pursuant to the RIFA for at least twelve months from the issuance date of these condensed consolidated financial statements. If we have not received full FDA approval and generated sufficient cash from product sales of YUTREPIA or are unable to access additional capital by the date of issuance of our fiscal year 2024 consolidated financial statements, we expect there would be substantial doubt about our ability to continue as a going concern as of that date. 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, which would have a material impact on our operations.â€² Basis of Presentation The audited interim condensed consolidated financial statements as of September 30, 2024 and for the three and nine months ended September 30, 2024 and 2023 have been prepared pursuant to the rules and regulations of the Securities and Exchange Commission (the â€œSECâ€) for interim financial reporting. These condensed consolidated financial statements are unaudited and, in the opinion of management, include all adjustments (consisting only of normal recurring adjustments and accruals) necessary for a fair statement of the results for the periods presented in accordance with accounting principles generally accepted in the United States of America (â€œGAAPâ€). The year-end condensed consolidated balance sheet data was derived from our audited consolidated financial statements but does not include all disclosures required by GAAP. Operating results for the three and nine months ended September 30, 2024 are not necessarily indicative of the results that may be expected for the fiscal year ending December 31, 2024. Certain information and footnote disclosures normally included in the annual consolidated financial statements prepared in accordance with GAAP have been omitted in accordance with the SECâ€™s rules and regulations for interim reporting. Certain amounts have been reclassified from the prior year presentation to conform to current presentation, specifically in relation to the balance sheet presentation of finance and operating leases. Our financial position, results of operations and cash flows are presented in U.S. Dollars. The accompanying unaudited condensed consolidated financial statements and related notes should be read in conjunction with our audited consolidated financial statements for the year ended December 31, 2023, which are included in our 2023 Annual Report on Form 10-K for the fiscal year ended December 31, 2023 (the â€œ2023 Annual Report on Form 10-Kâ€). Use of Estimates The preparation of financial statements in accordance with GAAP requires management to make estimates and assumptions that affect the reported amounts of assets and liabilities and the disclosure of contingent assets and liabilities at the date of the financial statements, as well as the reported amounts of revenues and expenses during the period. These estimates are based on historical experience and various other assumptions believed to be reasonable under the circumstances. We evaluate our estimates on an ongoing basis, including those related to the valuation of stock-based awards, certain accruals, the revenue interest financing payable, and intangible and contract acquisition cost amortization, and make changes to the estimates and related disclosures as our experience develops or new information becomes known. Actual results will most likely differ from those estimates. Segment Information GAAP requires segmentation based on an entityâ€™s internal organization and reporting of revenue and operating income based upon internal accounting methods commonly referred to as the â€œmanagement approach.â€ Operating segments are defined as components of an enterprise about which separate financial information is available that is evaluated regularly by the chief operating decision maker (CODM), or decision-making group, in deciding how to allocate resources and in assessing performance. Our CODM is our Chief Executive Officer. We have determined that we have one operating and reporting segment.â€¹² Table of Contents Summary of Significant Accounting Policies Our significant accounting policies are disclosed in Note 2 of the consolidated financial statements for the years ended December 31, 2023 and 2022, which are included in our 2023 Annual Report on Form 10-K. There have been no material changes to our significant accounting policies during the nine months ended September 30, 2024. Recent Accounting Pronouncements From time to time, new accounting pronouncements are issued by the Financial Accounting Standards Board under its accounting standards codifications (ASC) or other standard setting bodies and are adopted by us as of the specified effective date. For the nine months ended September 30, 2024, there were no newly adopted accounting pronouncements that had a material impact on our condensed consolidated financial statements. As of September 30, 2024, there are no recently issued but not yet adopted accounting pronouncements that are expected to materially impact our condensed consolidated financial statements. Cash, Cash Equivalents, and Concentration of Credit Risk We consider all highly liquid investments with a maturity of three months or less at the date of purchase to be cash equivalents. Financial instruments that potentially subject us to concentrations of credit risk consist of cash and cash equivalents. We are exposed to credit risk, subject to federal deposit insurance, in the event of default by the financial institutions holding our cash and cash equivalents to the extent of amounts recorded on the condensed consolidated balance sheet. Our cash and cash equivalents are held at multiple accredited financial institutions. We have not experienced any losses on such accounts and do not believe that we are subject to unusual credit risk beyond the normal credit risk associated with commercial banking relationships. Such deposits have exceeded and will continue to exceed federally insured limits. Accounts Receivable Accounts receivable are stated at net realizable value and net of an allowance for credit losses as of each balance sheet date, if applicable. One customer accounted for 96% and 99% of our accounts receivable, net at September 30, 2024 and December 31, 2023, respectively. As of September 30, 2024 and December 31, 2023, we have not recorded an allowance for credit losses. Prelaunch Inventory We capitalize prelaunch inventory prior to receiving regulatory approval if regulatory approval and subsequent commercialization of a product is probable and we also expect future economic benefit from the sales of the product to be realized. Prior to this conclusion, we expense prelaunch inventory as research and development expense in the period incurred. For prelaunch inventory that is capitalized, we consider a number of specific facts and circumstances, including the productâ€™s shelf life, the product's current status in the development and regulatory approval process, results from related clinical trials, results from meetings with relevant regulatory agencies prior to the filing of regulatory applications, potential obstacles to the approval process, viability of commercialization and market trends. In late 2023, based on our assessment of the legal and regulatory process related to YUTREPIA, we concluded that we met the criteria to capitalize expenditures for prelaunch inventory. We capitalized \$7.4 million of prelaunch inventory as of September 30, 2024 and none as of December 31, 2023. If either regulatory approval or market acceptance post-approval of YUTREPIA do not occur at all or on a timely basis prior to the inventory shelf-life expiration, we may be required to write-off some or all prelaunch inventory, which could affect our financial condition and financial results. Long-Lived Assets We review long-lived assets, including definite-life intangible assets, for realizability on an ongoing basis. Changes in depreciation and amortization, generally accelerated depreciation and variable amortization, are determined and recorded when estimates of the remaining useful lives or residual values of long-term assets change. We also review 13 Table of Contents for impairment when conditions exist that indicate the carrying amount of the assets may not be fully recoverable. In those circumstances, we perform undiscounted operating cash flow analyses to determine if an impairment exists. When testing for asset impairment, we group assets and liabilities at the lowest level for which cash flows are separately identifiable. Any impairment loss is calculated as the excess of the assetâ€™s carrying value over its estimated fair value. Fair value is estimated based on the discounted cash flows for the asset group over the remaining useful life or based on the expected cash proceeds for the asset less costs of disposal. Any impairment losses would be recorded in the consolidated statements of operations. To date, no such impairments have occurred. Goodwill We assess goodwill for impairment at least annually as of July 1 or whenever events or changes in circumstances indicate that the carrying amount of such assets may not be recoverable. For example, significant and unanticipated changes or our inability to obtain or maintain regulatory approvals for our product candidates, including the NDA for YUTREPIA, could trigger testing of our goodwill for impairment at an interim date. We have one reporting unit. We have the option to first assess qualitative factors to determine whether events or circumstances indicate it is more likely than not that the fair value of a reporting unit is greater than its carrying amount, in which case a quantitative impairment test is not required. Per ASC 350, Intangibles Goodwill and Other, the quantitative goodwill impairment test is performed by comparing the fair value of the reporting unit with its carrying amount, including goodwill. If the fair value of the reporting unit exceeds its carrying amount, goodwill is not impaired. An impairment loss is recognized for any excess of the carrying amount of the reporting unitâ€™s goodwill over the fair value up to the amount of goodwill allocated to the reporting unit. Income tax effects from any tax-deductible goodwill on the carrying amount of the reporting unit are considered when measuring the goodwill impairment loss, if applicable. As of September 30, 2024, we concluded there were no events or changes in circumstances which indicated that the carrying amount of goodwill was not recoverable. We completed our annual impairment test as of July 1, 2024 and concluded that no impairments had occurred. Revenue Interest Financing Payable We recognized a liability related to amounts received in January 2023, July 2023, and January 2024 pursuant to the RIFA under ASC 470-10, Debt and ASC 835-30, Interest - Imputation of Interest. The liability will be accreted under the effective interest method based upon the estimated amount of future payments to be made pursuant to the RIFA. If the timing or amounts of any estimated future payments change, we will prospectively adjust the effective interest and the related amortization of the liability and related issuance costs. A significant increase or decrease in these estimates could materially impact the liability balance and related interest expense. Amendments are assessed under ASC 470 to determine appropriate treatment as troubled debt restructurings, extinguishments or modifications. Revenue Recognition We recognize revenue in accordance with ASC 606, Revenue from Contracts with Customers (â€œASC 606â€). The core principle of ASC 606 is that a company should recognize revenue to depict the transfer of promised goods or services to customers in an amount that reflects the consideration to which the company expects to be entitled in exchange for those goods or services. The following five steps are applied to achieve that core principle:â€¹³ Step 1: Identify the contract with the customerâ€¹⁴ Step 2: Identify the performance obligations in the contractâ€¹⁵ Step 3: Determine the transaction priceâ€¹⁶ Step 4: Allocate the transaction price to the performance obligations in the contractâ€¹⁷ Step 5: Recognize revenue when the company satisfies a performance obligation In order to identify the performance obligations in a contract with a customer, we assess the promised goods or services in the contract and identify each promised good or service that is distinct. If a good or service is not distinct, the good or service is combined with other promised goods or services until a bundle of goods or services is identified that is distinct. The transaction price is the amount of consideration to which an entity expects to be entitled in exchange for transferring promised goods or services to a customer. The consideration promised in a contract with a customer may include fixed amounts, variable amounts, or both. Variable consideration is included in the transaction price only to the extent that it is probable that a significant reversal in the amount of cumulative revenue recognized will not occur when the uncertainty associated with the variable consideration is subsequently resolved. We evaluate any non-cash consideration, consideration payable to the customer, potential returns and refunds, and whether consideration contains a significant financing element in determining the transaction price. Revenue is measured based on consideration specified in a contract with a customer. We recognize revenue when it satisfies a performance obligation by transferring control over a service to a customer. The amount of revenue recognized reflects estimates for refunds and returns, which are presented as a reduction of accounts receivable where the right of setoff exists. Research and Development Expense Research and development costs are expensed as incurred in accordance with ASC 730, Research and Development and include facility-related costs related to research and development activities, direct costs from third parties, such as contract research organizations (â€œCROsâ€), contract

manufacturing organizations (CMOs), and consultants, as well as employee-related expenses, including salaries, benefits, and stock-based compensation. Research and development expenses also include costs of acquired product licenses and related technology rights where there is no alternative future use. Accrued Research and Development Expenses As part of the process of preparing the condensed consolidated financial statements, we are required to estimate accrued expenses. This process involves reviewing quotations and contracts, identifying services that have been performed on our behalf and estimating the level of service performed and the associated cost incurred for the service when we have not yet been invoiced or otherwise notified of the actual cost. The majority of our service providers invoice us monthly in arrears for services performed or when contractual milestones are met. We make estimates of our accrued expenses as of each balance sheet date in our condensed consolidated financial statements based on facts and circumstances known to us at that time. We periodically confirm the accuracy of these estimates with the service providers and make adjustments if necessary. The significant estimates in our accrued research and development expenses are related to expenses incurred with respect to CROs, CMOs and other vendors in connection with research and development and manufacturing activities. The financial terms of our agreements with CROs and CMOs are subject to negotiation, vary from contract to contract and may result in uneven payment flows. There may be instances in which payments made to our vendors will exceed the level of services provided and result in a prepayment of the applicable research and development or manufacturing expense. In accruing service fees, we estimate the time period over which services will be performed and the level of effort to be expended in each period. If the actual timing of the performance of services or the level of effort varies from such estimate, we adjust the accrual or prepaid expense accordingly. Although we do not expect our estimates to be materially different from amounts actually incurred, our understanding of the status and timing of services performed relative to the actual status and timing of services performed may vary and could result in us reporting amounts that are too high or too low in any particular period. There have been no material changes in estimates for the periods presented.

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Stock-Based Compensation We estimate the grant date fair value of stock-based awards and amortize this fair value to compensation expense over the requisite service period or the vesting period of the respective award. In arriving at stock-based compensation expense, we estimate the number of stock-based awards that will be forfeited due to employee turnover. The forfeiture assumption is based primarily on turn-over historical experience. If the actual forfeiture rate is higher than the estimated forfeiture rate, then an adjustment will be made to increase the estimated forfeiture rate, which will result in a decrease to the expense recognized in our financial statements. If the actual forfeiture rate is lower than the estimated forfeiture rate, then an adjustment will be made to lower the estimated forfeiture rate, which will result in an increase to expense recognized in our financial statements. The expense we recognize in future periods will be affected by changes in the estimated forfeiture rate and may differ from amounts recognized in the current period. See Note 9.

Net Loss Per Share Basic net loss per share is calculated by dividing net loss attributable to common stockholders by the weighted average shares outstanding during the period, without consideration of common stock equivalents. Diluted net loss per share is calculated by adjusting weighted average shares outstanding for the dilutive effect of common stock equivalents outstanding for the period, determined using the treasury-stock method. Due to their anti-dilutive effect, the calculation of diluted net loss per share excludes the following common stock equivalent shares: $\frac{\$450,000}{\$450,000}$ Total $\frac{\$12,695,261}{\$12,695,261}$ $\frac{\$11,728,033}{\$11,728,033}$ $\frac{\$12,885,288}{\$12,885,288}$ $\frac{\$11,634,831}{\$11,634,831}$. Certain common stock warrants are included in the calculation of basic and diluted net loss per share since their exercise price is de minimis. Fair Value Measurements ASC 825, Financial Instruments defines fair value as the amount that would be received to sell an asset or paid to transfer a liability in an orderly transaction between market participants (an exit price). As such, fair value is a market-based measurement that should be determined based on assumptions that market participants would use in pricing an asset or liability. ASC 825 establishes a three-tiered approach for valuation of financial instruments, which requires that fair value measurements be classified and disclosed in one of three tiers, whether or not recognized on our condensed consolidated balance sheets at fair value. The fair value hierarchy defines three-level valuation hierarchy for disclosure of fair value measurements as follows: Level 1 $\frac{\$7,912}{\$7,912}$ Quoted prices in active markets for identical assets or liabilities; Level 2 $\frac{\$7,912}{\$7,912}$ Inputs other than quoted prices included in active markets that are observable for the asset or liability, either directly or indirectly; and Level 3 $\frac{\$7,912}{\$7,912}$ Unobservable inputs for the asset and liability used to measure fair value, to the extent that observable inputs are not available. The categorization of a financial instrument within the valuation hierarchy is based upon the lowest level of input that is significant to the fair value measurement. The following table presents the placement in the fair value hierarchy of financial assets and liabilities measured at fair value as of September 30, 2024 and December 31, 2023.

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Quoted and Significant $\frac{\$7,912}{\$7,912}$ Quoted prices in active markets for identical assets or liabilities; $\frac{\$7,912}{\$7,912}$ Inputs other than quoted prices included in active markets that are observable for the asset or liability, either directly or indirectly; and $\frac{\$7,912}{\$7,912}$ Unobservable inputs for the asset and liability used to measure fair value, to the extent that observable inputs are not available. The categorization of a financial instrument within the valuation hierarchy is based upon the lowest level of input that is significant to the fair value measurement. The following table presents the placement in the fair value hierarchy of financial assets and liabilities measured at fair value as of September 30, 2024 and December 31, 2023.

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Market Funds $\frac{\$7,912}{\$7,912}$ Quoted prices in active markets for identical assets or liabilities; $\frac{\$7,912}{\$7,912}$ Inputs other than quoted prices included in active markets that are observable for the asset or liability, either directly or indirectly; and $\frac{\$7,912}{\$7,912}$ Unobservable inputs for the asset and liability used to measure fair value, to the extent that observable inputs are not available. The categorization of a financial instrument within the valuation hierarchy is based upon the lowest level of input that is significant to the fair value measurement. The following table presents the placement in the fair value hierarchy of financial assets and liabilities measured at fair value as of September 30, 2024 and December 31, 2023.

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Raw Materials $\frac{\$2,607}{\$2,607}$ Work in process; $\frac{\$4,835}{\$4,835}$ Inventory; $\frac{\$7,442}{\$7,442}$ Recognized as: $\frac{\$7,404}{\$7,404}$ Inventories; $\frac{\$384}{\$384}$ Other assets; $\frac{\$7,404}{\$7,404}$ Pre-launch Inventory; $\frac{\$7,912}{\$7,912}$ Capitalized costs of \$7.4 million associated with the production of YUTREPIA as a result of our determination that regulatory approval and subsequent commercialization is probable, and we also expect future economic benefit from the sales of YUTREPIA to be realized. Amounts recognized as Other Assets are comprised entirely of raw materials and work in process inventories not expected to be sold within one year of the balance sheet date.

Property, Plant and Equipment $\frac{\$6,953}{\$6,953}$ Office equipment; $\frac{\$6,834}{\$6,834}$ Furniture and fixtures; $\frac{\$19}{\$19}$ Computer and other equipment; $\frac{\$740}{\$740}$ Leasehold improvements; $\frac{\$11,450}{\$11,450}$ Construction in progress; $\frac{\$4,315}{\$4,315}$ Total property, plant and equipment.

Intangible Assets $\frac{\$7,784}{\$7,784}$ Net intangible assets; $\frac{\$4,480}{\$4,480}$ We recorded depreciation and amortization expense related to property, plant and equipment of \$0.3 million and \$0.3 million for the three months ended September 30, 2024 and 2023, respectively, and of \$0.8 million and \$0.9 million for the nine months ended September 30, 2024 and 2023, respectively.

Contract Acquisition Costs and Intangible Asset Contract acquisition costs and intangible asset are summarized as follows: $\frac{\$12,980}{\$12,980}$ Net Carrying Amount; $\frac{\$5,547}{\$5,547}$ Accumulated Amortization; $\frac{\$7,433}{\$7,433}$ Net Carrying Amount.

Contract Acquisition Costs $\frac{\$12,980}{\$12,980}$ Net Carrying Amount; $\frac{\$5,547}{\$5,547}$ Accumulated Amortization; $\frac{\$7,433}{\$7,433}$ Net Carrying Amount.

Intangible Asset $\frac{\$5,058}{\$5,058}$ Net Carrying Amount; $\frac{\$7,922}{\$7,922}$ Intangible asset; $\frac{\$(2,401)}{\$(2,401)}$ Accrued other expenses.

Accrued Other Expenses $\frac{\$3,219}{\$3,219}$ Accrued other expenses.

Stockholders' Equity $\frac{\$5,620}{\$5,620}$ Common Stock.

Common Stock $\frac{\$2,190}{\$2,190}$ Common Stock.

Accrued Research and Development Expenses $\frac{\$17,039}{\$17,039}$ Accrued research and development expenses.

Stockholders' Equity $\frac{\$13,400}{\$13,400}$ Stockholders' equity.

Common Stock $\frac{\$48}{\$48}$ Common Stock.

Accrued Research and Development Expenses $\frac{\$2,902}{\$2,902}$ Accrued research and development expenses.

Stockholders' Equity $\frac{\$1,916}{\$1,916}$ Stockholders' equity.

Accrued Other Expenses $\frac{\$4,254}{\$4,254}$ Accrued other expenses.

Accrued Research and Development Expenses $\frac{\$1,954}{\$1,954}$ Accrued research and development expenses.

Total Accrued Expenses and Other Current Liabilities $\frac{\$17,039}{\$17,039}$ Total accrued expenses and other current liabilities.

Stockholders' Equity $\frac{\$1,954}{\$1,954}$ Stockholders' equity.

Common Stock $\frac{\$1,954}{\$1,954}$ Common Stock.

Accrued Research and Development Expenses $\frac{\$17,039}{\$17,039}$ Accrued research and development expenses.

Stockholders' Equity $\frac{\$1,954}{\$1,954}$ Stockholders' equity.

Common Stock $\frac{\$1,954}{\$1,954}$ Common Stock.

Accrued Research and Development Expenses $\frac{\$17,039}{\$17,039}$ Accrued research and development expenses.

Stockholders' Equity $\frac{\$1,954}{\$1,954}$ Stockholders' equity.

Common Stock $\frac{\$1,954}{\$1,954}$ Common Stock.

Accrued Research and Development Expenses $\frac{\$17,039}{\$17,039}$ Accrued research and development expenses.

Stockholders' Equity $\frac{\$1,954}{\$1,954}$ Stockholders' equity.

Common Stock $\frac{\$1,954}{\$1,954}$ Common Stock.

Accrued Research and Development Expenses $\frac{\$17,039}{\$17,039}$ Accrued research and development expenses.

Stockholders' Equity $\frac{\$1,954}{\$1,954}$ Stockholders' equity.

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current conditions as set forth in the Pharmosa License Agreement. Mainbridge Health Care Device Development and Supply Agreement In December 2022, we entered into a Device Development and Supply Agreement (the "Pump Development Agreement") with Mainbridge Health Partners, LLC ("Mainbridge") and Sandoz Inc. ("Sandoz"). The Pump Development Agreement provides for the cooperation between us, Sandoz and Mainbridge to develop a new pump that is suitable for the subcutaneous administration of Treprostин Injection. Mainbridge will perform all development, validation and testing activities required for the pump and related consumables in anticipation of submitting a 510(k) clearance application for the pump to the FDA. In connection with the Pump Development Agreement, we and Sandoz have agreed to pay Mainbridge certain future contingent milestone payments in accordance with the terms and conditions set forth therein. UNC License Agreement We perform research under a license agreement with The University of North Carolina at Chapel Hill ("UNC") as amended to date (the "UNC License Agreement"). As part of the UNC License Agreement, we hold an exclusive license to certain research and development technologies and processes in various stages of patent pursuit, for use in our research and development and commercial activities, with a term until the expiration date of the last to expire patent subject to the UNC License Agreement, subject to industry standard contractual compliance. Under the UNC License Agreement, we are obligated to pay UNC royalties equal to a low single digit percentage of all net sales of drug products whose manufacture, use or sale includes any use of the technology or patent rights covered by the UNC License 28Table of Contents Agreement, including YUTREPIA. We may grant sublicenses of UNC licensed intellectual property in return for specified payments based on a percentage of any fee, royalty or other consideration received. Chasm Technologies In March 2012, we entered into an agreement, as amended, with Chasm Technologies, Inc. for manufacturing consulting services related to our manufacturing capabilities during the term of the agreement. We agreed to pay future contingent milestones and royalties on net sales totaling no more than \$1.5 million, \$0.2 million of which has been accrued as of September 30, 2024. Employment Agreements and Executive Severance and Change in Control Plan We have agreements with certain employees and an Executive Severance and Change in Control Plan which covers certain other employees which require payments if certain events, such as a change in control or termination without cause, occur. Purchase Obligations We enter into contracts in the normal course of business with contract service providers to assist in the performance of research and development and manufacturing activities. Subject to required notice periods and obligations under binding purchase orders, we can elect to discontinue the work under these agreements at any time. On July 14, 2023, we entered into an Amended and Restated Commercial Manufacturing Services and Supply Agreement with Lonza Tampa LLC ("Lonza") (the "CSA"). Lonza is our sole supplier for encapsulation and packaging services for YUTREPIA. Pursuant to the terms of the CSA, we deliver bulk treprostин powder, manufactured using our proprietary PRINTA® technology, and Lonza encapsulates and packages it. The CSA was effective upon signing and will be in effect for an initial term of 5 years from receipt of regulatory approval of YUTREPIA by the FDA ("Regulatory Approval") absent termination by either party in accordance with the terms of the CSA. We may terminate the CSA upon 60 days' written notice to Lonza in the event that the application for regulatory approval is rejected by the FDA and such FDA decision is not caused by the fault of the Company (the "Termination for FDA Rejection"). Lonza may terminate the CSA upon 120 days written notice if we do not receive regulatory approval by December 31, 2024 (the "Termination for FDA Delay"). Upon any Termination for FDA Rejection or Termination for FDA Delay, we would reimburse Lonza for 50% of its documented out-of-pocket expenditures for any capital equipment that is purchased by Lonza after the effective date of the Agreement to perform the services for us, not to exceed \$2.5 million in the aggregate. We are required to provide Lonza with quarterly forecasts of our expected production requirements for the following 24-month period, the first twelve months of which is considered a binding, firm order. We are required to purchase certain minimum annual order quantities, which may be adjusted by us after the thirteenth month after receipt of regulatory approval (as defined in the CSA). The CSA provides for tiered pricing depending upon the batch size ordered. In addition, we are party to a multi-year supply agreement with LGM Pharma, LLC (LGM) to supply active pharmaceutical ingredients for YUTREPIA. Under the supply agreement with LGM, we are required to provide rolling forecasts, a portion of which will be considered a binding, firm order, subject to an annual minimum purchase commitment of \$2.7 million for the term of the agreement. The agreement expires five years from the first marketing authorization approval of YUTREPIA. As of September 30, 2024, we have non-cancelable commitments for product manufacturing and supply costs of approximately \$14.7 million. Other Contingencies and Commitments From time-to-time we are subject to claims and litigation in the normal course of business, none of which do we believe represent a risk of material loss or exposure. See Note 14 for further discussion of pending legal proceedings. 29Table of Contents In addition to the commitments described above, we are party to other commitments, including non-cancelable leases and long-term debt, which are described elsewhere in these notes to the condensed consolidated financial statements. 14. Legal Proceedings YUTREPIA-Related Litigation In 2020, United Therapeutics filed a complaint for patent infringement against the Company in the U.S. District Court for the District of Delaware (the "Original Hatch-Waxman Litigation"), asserting infringement by the Company of U.S. Patent Nos. 9,604,901, entitled "Process to Prepare Treprostин, the Active Ingredient in Remodulin®" (the "901 Patent"), 9,593,066, entitled "Process to Prepare Treprostин, the Active Ingredient in Remodulin®" (the "906 Patent"), and 10,716,793, entitled "Treprostин Administration by Inhalation" (the "793 Patent") relating to United Therapeutics' Tyvaso® (a nebulized treprostин solution for the treatment of PAH). United Therapeutics' complaint was in response to the Company's NDA for YUTREPIA, filed with the FDA, requesting approval to market YUTREPIA, a dry powder formulation of treprostин for the treatment of PAH. The YUTREPIA NDA was filed under the 505(b)(2) regulatory pathway with Tyvaso® as the reference listed drug. In December 2021, United Therapeutics filed a stipulation of partial judgment with respect to the "901 Patent under which United Therapeutics agreed to the entry of judgment of the Company's non-infringement of the "901 Patent. In August 2022, Judge Andrews, who was presiding over the Original Hatch-Waxman Litigation, issued an opinion that claims 1, 2, 3, 6 and 9 of the "906 Patent were invalid, that the remaining asserted claims of the "906 Patent were not infringed by the Company. In January 2021, the Company filed a petition for inter partes review with the Patent Trial and Appeal Board (the "PTAB") relating to the "793 Patent, seeking a determination that the claims in the "793 Patent are invalid. In July 2022, the PTAB ruled in the Company's favor, concluding that based on the preponderance of the evidence, all the claims of the "793 Patent have been shown to be unpatentable. These decisions related to the "901 Patent, the "906 Patent and the "793 Patent have all been affirmed on appeal and are not subject to further appeal. In connection with an amendment to the Company's NDA filed in July 2023 to add PH-ILD as an indication for YUTREPIA, the Company provided a new notice of the paragraph IV certification to United Therapeutics as the owner of the patents that are the subject of the certification to which the NDA for YUTREPIA refers. As a result, in September 2023, United Therapeutics filed a second complaint for patent infringement against the Company in the U.S. District Court for the District of Delaware (Case No. 1:23-cv-00975-RGA) (the "New Hatch-Waxman Litigation"), again asserting infringement by the Company of the "793 Patent. In November 2023, the U.S. Patent and Trademark Office (the "USPTO") issued U.S. Patent No. 11,826,327, or the "327 Patent, entitled "Treatment for Interstitial Lung Disease", to United Therapeutics. On November 30, 2023, United Therapeutics filed an amended complaint in the New Hatch-Waxman Litigation asserting infringement of the "327 Patent by the practice of YUTREPIA based on the amended NDA. In January 2024, the Company filed an answer, counterclaims and a partial motion to dismiss the claims related to the "793 Patent as a result of the decision by the United States Court of Appeals for the Federal Circuit to affirm the PTAB's finding that the "793 patent is unpatentable. In February 2024, United Therapeutics stipulated to the dismissal of the claims in the New Hatch-Waxman Litigation related to the "793 Patent. In February 2024, United Therapeutics also filed a motion seeking a preliminary injunction to prevent the Company from manufacturing, marketing, storing, importing, distributing, offering for sale, and/or selling YUTREPIA for the treatment of PH-ILD. Judge Andrews denied the motion for a preliminary injunction in May 2024. Discovery in the case remains ongoing. FDA Litigation In February 2024, United Therapeutics filed a complaint against the FDA in the U.S. District Court for the District of Columbia (the "D.C. District Court"), challenging the FDA's acceptance of the Company's amended NDA for review (the "Original FDA Litigation"). The Company intervened and became a party to the lawsuit in March 2024. In March 2024, United Therapeutics filed a motion for a temporary restraining order and preliminary injunction in the FDA Litigation, seeking to enjoin the FDA from approving the Company's NDA for YUTREPIA with respect to the indication to treat PH-ILD. United Therapeutics' motion was denied in March 2024. On August 20, 2024, United Therapeutics voluntarily dismissed its complaint, without prejudice. 30Table of Contents On August 21, 2024, the Company filed a lawsuit in the D.C. District Court to challenge the decision by the FDA to grant three-year regulatory exclusivity to Tyvaso DPI (the "New FDA Litigation"). The D.C. District Court has granted the parties' motion for an expedited summary judgment briefing schedule, with briefing to be completed on or before November 15, 2024 and a summary judgment hearing scheduled for December 5, 2024. On September 16, 2024, United Therapeutics filed a cross claim in the New FDA Litigation, re-asserting its challenge to FDA's acceptance of the Company's amended NDA for review. The Company intervened and became a party with respect to the cross claim on November 5, 2024. United Therapeutics has filed a motion for an expedited summary judgment briefing schedule for its cross claim. That motion remains pending. Trade Secret Litigation In December 2021, United Therapeutics filed a complaint in the Superior Court in Durham County, North Carolina, alleging that the Company and Robert Roscigno (the "Dr. Roscigno") a former United Therapeutics employee, who later joined the Company as an employee many years after terminating his employment with United Therapeutics, conspired to misappropriate certain trade secrets of United Therapeutics and engaged in unfair or deceptive trade practices. In January 2024, Dr. Roscigno filed a motion for summary judgment with respect to all claims, but the motion was denied in July 2024. In addition, in July 2024, the Company filed a motion for summary judgment with respect to all claims. A hearing on the Company's motion for summary judgment has been scheduled for December 19, 2024. In May 2024, United Therapeutics filed a second complaint in the Superior Court in Durham County, North Carolina, against Dr. Roscigno, alleging that he breached prior employment agreements with United Therapeutics by failing to assign to United Therapeutics his interest in patents obtained by the Company that relied upon or benefited from certain inventions, discoveries, materials, authorship, derivatives and results developed by Dr. Roscigno while he was employed by United Therapeutics. The Company was also named as a defendant in this new lawsuit. As part of the lawsuit, United Therapeutics alleges that Dr. Roscigno misappropriated certain intellectual property of United Therapeutics which led to the development of YUTREPIA. The complaint also seeks declaratory judgement such that all right, title and interest in and to any patentable or unpatentable inventions, discoveries, and ideas made or conceived by Dr. Roscigno while employed by the Company should be assigned and transferred to United Therapeutics because they involved the use of United Therapeutics' confidential information. On July 30, 2024, the Company filed a motion to dismiss all claims. A hearing on the Company's motion to dismiss has been scheduled for December 19, 2024. 4. RareGen Litigation In April 2019, Sandoz and Liquidia PAH (then known as RareGen) filed a complaint against United Therapeutics and Smiths Medical (now ICU Medical) in the District Court of New Jersey (Case No. No. 3:19-cv-10170), (the "RareGen Litigation"), alleging that United Therapeutics and Smiths Medical violated the Sherman Antitrust Act of 1890, state law antitrust statutes and unfair competition statutes by engaging in anticompetitive acts regarding the drug treprostин for the treatment of PAH. In March 2020, Sandoz and Liquidia PAH filed a first amended complaint adding a claim that United Therapeutics breached a settlement agreement that was entered into in 2015, in which United Therapeutics agreed to not interfere with Sandoz's efforts to launch its generic treprostин, by taking calculated steps to restrict and interfere with the launch of Sandoz's competing generic product. United Therapeutics developed treprostин under the brand name Remodulin® and Smiths Medical manufactured a pump and cartridges that are used to inject treprostин into patients continuously throughout the day. Sandoz and Liquidia PAH allege that United Therapeutics and Smiths Medical entered into anticompetitive agreements (i) whereby Smiths Medical placed restrictions on the cartridges such that they can only be used with United Therapeutics' branded Remodulin® product and (ii) requiring Smiths Medical to enter into agreements with specialty pharmacies to sell the cartridges only for use with Remodulin®. In November 2020, Sandoz and Liquidia PAH entered into a binding term sheet (the "Term Sheet") with Smiths Medical in order to resolve the outstanding RareGen Litigation solely with respect to disputes between Smiths Medical, Liquidia PAH and Sandoz. In April 2021, Liquidia PAH and Sandoz entered into a Long Form Settlement Agreement (the "Settlement Agreement") with Smiths Medical to further detail the terms of the settlement among such parties as reflected in the Term Sheet. Pursuant to the Term Sheet and the Settlement Agreement, the former RareGen members and Sandoz received a payment of \$4.25 million that was evenly split between the parties. In addition, pursuant to the Term Sheet and Settlement Agreement, Smiths Medical disclosed and made available to Sandoz and Liquidia PAH 31Table of Contents certain specifications and other information related to the cartridge that Smiths Medical developed and manufactures for use with the CADD-MS 3 infusion pump (the "CADD-MS 3 Cartridge"). Pursuant to the Settlement Agreement, Smiths Medical also granted Liquidia PAH and Sandoz a non-exclusive, royalty-free license in the United States to Smiths Medical's patents and copyrights associated with the CADD-MS 3 Cartridge and certain other information for use of the CADD-MS 3 pump and the CADD-MS 3 Cartridges. Smiths also agreed in the Settlement Agreement to provide information and assistance in support of Liquidia PAH's efforts to receive FDA clearance for the RG 3ml Medication Cartridge (the "RG Cartridge") and to continue to service certain CADD-MS 3 pumps that are available for use with the Treprostин Injection through January 1, 2025. Liquidia PAH and Sandoz agreed, among other things, to indemnify Smiths from certain liabilities related to the RG Cartridge. In September 2021, United Therapeutics filed a motion for summary judgment with respect to all of the claims brought by Sandoz and Liquidia PAH against United Therapeutics. At the same time, Sandoz filed a motion for summary judgment with respect to the breach of contract claim. In March 2022, the Court issued an order granting partial summary judgment to United Therapeutics with respect to the antitrust and unfair competition claims, denying summary judgment to United Therapeutics with respect to the breach of contract claim, and granting partial summary judgment to Sandoz with respect to the breach of contract claim. A trial to determine the amount of damages due from United Therapeutics to Sandoz with respect to the breach of contract claim was held from late April to early May 2024, and closing arguments were held in June 2024. On November 1, 2024, the Court entered a judgment in the amount of \$70.6 million. United Therapeutics has filed a notice of its intent to appeal the decision. Under the Promotion Agreement, all proceeds from the litigation will be divided evenly between Sandoz and Liquidia PAH. Under the litigation finance agreements that Liquidia PAH has entered into with Henderson and PBM, any net proceeds received by Liquidia PAH with respect to the RareGen Litigation will be divided between Henderson and PBM. 4.1 First Amendment to the Pharmosa License Agreement (the "First Amendment") and concurrently entered into a Device License Agreement with Pharmosa. See Note 13 for additional information. 4.2 Management's Discussion and Analysis of Financial Condition and Results of Operations. You should read the following discussion and analysis of our financial condition and results of operations together with our condensed consolidated financial statements and related notes appearing in this Quarterly Report on Form 10-Q. This discussion and other parts of this Quarterly Report contain forward-looking statements that involve risks and uncertainties, such as statements of our plans, objectives, expectations and intentions. As a result of many factors, including those factors set forth in the "Risk Factors" section of this Quarterly Report, our actual results could differ materially from the results described in, or implied by, the forward-looking statements contained in the following discussion and analysis. Objective The following Management's Discussion and Analysis of Financial Condition and Results of Operations is intended to provide information necessary to understand our condensed consolidated financial statements and highlight certain other information which, in the opinion of management, will enhance a reader's understanding of our financial condition, changes in financial condition and results of operations. In particular, the discussion is intended to provide an analysis of significant trends and material changes in our financial position and the operating results of our business during the three and nine months ended September 30, 2024 as compared to the three and nine months ended September 30, 2023. Also refer to our Annual Report on Form 10-K for the fiscal year ended December 31, 2023, which includes detailed discussions of various items impacting our business, results of operations and financial condition. 32Table of Contents Overview We are a biopharmaceutical company focused on the development, manufacture, and commercialization of products that address unmet patient needs, with current focus directed towards rare cardiopulmonary diseases such as pulmonary arterial hypertension ("PAH") and pulmonary hypertension associated with interstitial lung disease ("PH-ILD"). We operate through our wholly owned operating subsidiaries, Liquidia Technologies, Inc. ("Liquidia Technologies") and Liquidia PAH, LLC ("Liquidia PAH"), formerly known as RareGen, LLC ("RareGen"). We currently generate revenue pursuant to a promotion agreement between Liquidia PAH and Sandoz Inc. ("Sandoz"), dated as of August 1, 2018, as amended (the "Promotion Agreement"), sharing profit derived from the sale of Sandoz's substitutable generic treprostин injection ("Treprostин Injection") in the United States. Liquidia PAH has the exclusive rights to conduct commercial activities to encourage the appropriate use of Treprostин Injection. We employ a targeted sales force calling on physicians and hospital pharmacies involved in the treatment of PAH and PH-ILD in the United States, as well as key stakeholders involved in the distribution and reimbursement of medicines to treat these patients. We established our commercial presence in the field to support Treprostин Injection and have since expanded our presence to support the potential launch of YUTREPIA (treprostин) inhalation powder ("YUTREPIA"), further validating our reputation as a company committed to supporting PAH and PH-ILD patients. We conduct research, development and manufacturing of novel products by applying our subject matter expertise in cardiopulmonary

Financial Reporting There were no changes in our internal control over financial reporting during the quarter ended September 30, 2024 that have materially affected, or are reasonably likely to materially affect, our internal control over financial reporting. **Legal Proceedings** For information on our legal proceedings, see Note 14 to the condensed consolidated financial statements appearing elsewhere in this Quarterly Report on Form 10-Q. **Risk Factors** Investing in our common stock involves a high degree of risk. You should carefully consider the risks described below, as well as the other information in this Quarterly Report on Form 10-Q, including our financial statements and the related notes thereto. **Management's Discussion and Analysis of Financial Condition and Results of Operations** and the information contained under the heading "Cautionary Note Regarding Forward-Looking Statements" before deciding whether to invest in our common stock. The occurrence of any of the events or developments described below could harm our business, financial condition, results of operations and growth prospects. In such an event, the market price of our common stock could decline and you may lose all or part of your investment. Additional risks and uncertainties not presently known to us or that we currently deem immaterial also may impair our business operations. We may update these risk factors in our periodic and other filings with the SEC. The following is a summary of the principal risk factors described in this section: **—We expect to incur significant expenses and operating losses for the foreseeable future as we advance our product candidates through clinical trials, seek regulatory approval and pursue commercialization of any approved product candidates. The future viability of our company will depend on our ability to fund future operations and capital requirements with potential sales of any approved product candidates and/or with additional capital from external financing.** **—We have a history of losses and our future profitability remains uncertain.** **—We are primarily dependent on the success of our product candidates, YUTREPIA and L606, and these product candidates may fail to receive final marketing approval (in a timely manner or at all) for some or all of the indications for which we are seeking approval or may not be commercialized successfully.** **—United Therapeutics has initiated multiple lawsuits against us in which it has claimed that YUTREPIA is infringing its patents, a separate lawsuit against us that we and a former United Therapeutics employee, who later joined us as an employee, conspired to misappropriate certain trade secrets of United Therapeutics and engaged in unfair or deceptive trade practices, and a separate lawsuit against the FDA asserting that the FDA improperly accepted for review an amendment to our NDA for YUTREPIA.** These lawsuits, and other lawsuits that United Therapeutics may file in the future, may result in our company being further delayed in its efforts to commercialize YUTREPIA or result in substantial damage claims against us if we launch YUTREPIA and we are later found to infringe. **—Liquida PAH does not hold the FDA regulatory approval for Treprostinil Injection, the RG Cartridge or pumps used to administer Treprostinil Injection and is dependent on Sandoz, Chengdu and the pump manufacturers to manufacture and supply Treprostinil Injection, the RG Cartridge and pumps used to administer Treprostinil Injection, respectively, in compliance with FDA requirements, and is more broadly dependent on their FDA and healthcare compliance relative to Treprostinil Injection, the RG Cartridge and the pumps used to administer Treprostinil Injection, respectively.** **—Treprostinil Injection is presently administered subcutaneously via ICU Medical's CADD-MS 3 infusion pump. ICU Medical no longer manufactures the CADD-MS 3 infusion pump and has indicated its intention to discontinue service and maintenance of CADD-MS 3 infusion pumps after January 1, 2025. Should components of the CADD-MS 3 pump become unavailable, ICU Medical's ability to service and maintain such pumps may terminate earlier than anticipated.** For instance, during 2022 we became aware of a potential shortage of a critical component of the CADD-MS 3 infusion pump that may cause the number of CADD-MS 3 infusion pumps available for the administration of Treprostinil Injection to 46Table of Contentsbe depleted prior to January 1, 2025. In the event the specialty pharmacies are unable to access sufficient quantities of operable pumps or in the event we are unable to identify or develop a new pump prior to the current pumps becoming unavailable, the commercial success of Treprostinil Injection may be adversely affected. **—Sales of Treprostinil Injection are dependent on market acceptance of generic treprostinil for parenteral administration and the medical devices used for administration of Treprostinil Injection, including the ICU Medical infusion pumps, any future pumps that we develop, and the RG Cartridge, by patients, health care providers and by third-party payors, while interactions with these persons and entities are subject to compliance requirements.** The commercial success of Treprostinil Injection may also be impacted by increasing generic competition which may result in declining prices for Treprostinil Injection. **—We expect that we will need further financing for our existing business and future growth, which may not be available on acceptable terms, if at all.** Failure to obtain funding on acceptable terms and on a timely basis may require us to curtail, delay or discontinue our product development efforts or other operations. The failure to obtain further financing may also prevent us from capitalizing on other potential product candidates or indications which may be more profitable than YUTREPIA and/or L606 or for which there may be a greater likelihood of success. **—We face significant competition from large pharmaceutical companies, among others, in developing our products and in gaining regulatory approval to bring them to market in time to achieve commercial success, and our operating results will suffer if we are unable to compete effectively, including if one or more such products have a superior product profile to YUTREPIA and/or L606.** **—Our financing facility with HCR contains operating and financial covenants that restrict our business and financing activities, and is subject to acceleration in specified circumstances, which may result in HCR taking possession and disposing of any collateral.** **—Our products may not achieve market acceptance or third-party payor coverage.** **—Our product candidates are based on proprietary, novel technology, which have not been used to manufacture any products that have been previously approved by the FDA, making it difficult to predict the time and cost of development and of subsequently obtaining final regulatory approval.** In addition, we may experience unexpected challenges as we ramp up our manufacturing capacity to meet demand or during commercial manufacturing, which may result in our inability to supply sufficient quantities of product to meet demand. **—Our business and operations may be adversely affected by the effects of health epidemics.** **—We may not be able to build or maintain a commercial operation, including establishing and maintaining marketing and sales capabilities or entering into agreements with third parties to market and sell our drug products.** **—We depend on third parties for clinical and commercial supplies, including single suppliers for the active ingredient, the device, encapsulation and packaging of YUTREPIA and single suppliers for the drug product and device for L606.** In the event of any disruption in these suppliers, our ability to develop and commercialize, and the timeline for commercialization of, YUTREPIA and/or L606 may be adversely affected. **—We rely on third parties to conduct our preclinical studies and clinical trials.** **—We may become involved in litigation to protect our intellectual property, to enforce our intellectual property rights or to defend against claims of intellectual property infringement by third parties, which could be expensive, time-consuming and may not be successful.** **—We depend on skilled labor, and our business and prospects may be adversely affected if we lose the services of our skilled personnel, including those in senior management, or are unable to attract new skilled personnel.** **—We expect that the market price of our common stock may be volatile, and you may lose all or part of your investment.** **Risks Related to our Financial Position and Need for Additional Capital** We expect to incur significant expenses and operating losses for the foreseeable future as we advance our product candidates through clinical trials, seek regulatory approval and pursue commercialization of any approved product candidates. The future viability of our company will depend on our ability to fund future operations and capital requirements with potential sales of any approved product candidates and/or with additional capital from external financing. We are subject to risks and uncertainties common to early-stage companies in the biotechnology industry, including, but not limited to, development by competitors of new technological innovations, dependence on key personnel, protection of proprietary technology, compliance with government regulations, and the ability to secure additional capital to fund operations. We expect to incur significant expenses and may incur significant operating losses for the foreseeable future as we advance product candidates through clinical trials, seek regulatory approval and pursue commercialization of any approved product candidates. We do not expect to generate significant revenue unless and until we are able to obtain marketing approval for and successfully commercialize one or more of our product candidates. In addition, if we obtain marketing approval for any of our product candidates, we would incur significant commercialization expenses related to product manufacturing, marketing, sales and distribution. These efforts require significant amounts of capital, adequate personnel and infrastructure, and extensive compliance-reporting capabilities. If we have not received full FDA approval and generated sufficient cash from product sales of YUTREPIA or are unable to access additional capital by the date of issuance of our fiscal year 2024 consolidated financial statements, we expect there would be substantial doubt about our ability to continue as a going concern as of that date. **—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, which would have a material impact on our operations.** Even if our development efforts are successful, it is uncertain when, if ever, we will realize significant revenue from product sales. The future viability of our company will depend on our ability to fund future operations and capital requirements with potential sales of any approved product candidates and/or with additional capital from external financing. We may seek additional funding through public or private financings, debt financing or collaboration. Our inability to obtain funding, when needed, would have a negative impact on our financial condition and ability to pursue our business strategies. **—We have a history of losses and our future profitability remains uncertain.** We have incurred net losses of \$92.0 million during the nine months ended September 30, 2024, and \$78.5 million and \$41.0 million during the years ended December 31, 2023 and 2022, respectively. We also had negative operating cash flows for each of these periods. As of September 30, 2024, we had an accumulated deficit of \$521.1 million. Since our incorporation, we have invested heavily in the development of our product candidates and technologies, as well as in recruiting management and scientific personnel. To date, we have not commenced the commercialization of our product candidates and all of our revenue has been derived from up-front fees and milestone payments made to us in connection with licensing and collaboration arrangements we have entered into and the Promotion Agreement, under which we share in the profit derived from the sale of Treprostinil Injection in the United States. These up-front fees and milestone payments have been, and combined with revenue generated from Treprostinil Injection may continue to be, insufficient to match our operating expenses. We expect to continue to devote substantial financial and other resources to the clinical development of our product candidates and, as a result, must generate significant revenue to achieve and maintain profitability or we will need to raise additional capital to continue funding clinical development. We may continue to incur losses and negative cash flow and may never transition to profitability or positive cash flow. **—We expect that we will need further financing for our existing business and future growth, which may not be available on acceptable terms, if at all.** Failure to obtain funding on acceptable terms and on a timely basis may require us to curtail, delay or discontinue our product development efforts or other operations. The failure to obtain further financing may also prevent us from capitalizing on other potential product candidates or indications which may be more profitable than YUTREPIA and/or L606 or for which there may be a greater likelihood of success. We expect that we will need to raise additional funds to meet our future funding requirements for the continued research, development and commercialization of our product candidates and technology. Our future funding requirements will be heavily determined by the timing of the potential commercialization of YUTREPIA and the resources needed to support development of our product candidates. In the event that funds generated from our operations are insufficient to fund our future growth, we may raise additional funds through the issuance of equity or debt securities or by borrowing from banks or other financial institutions. We cannot assure you that we will be able to obtain such additional financing on terms that are acceptable to us, or at all. Global and local economic conditions could negatively affect our ability to raise funds. To the extent that we raise additional capital through the sale of equity or convertible debt securities, your ownership interest will be diluted, and the terms of such securities may include liquidation or other preferences that adversely affect your rights as a stockholder. Such financing, even if obtained, may be accompanied by restrictive covenants that may, among others, limit our ability to pay dividends or require us to seek consent for payment of dividends, or restrict our freedom to operate our business by requiring consent for certain actions. **—If we fail to obtain financing on terms that are favorable to us, we will not be able to implement our growth plans, and we may be required to significantly curtail, delay or discontinue one or more of our research, development or manufacturing programs or the commercialization of any approved product.** Furthermore, if we fail to obtain additional financing on terms that are acceptable to us, we may forgo or delay the pursuit of opportunities presented by other potential product candidates or indications that may later prove to have greater commercial potential than the product candidates and indications that we have chosen to pursue. **—Our financing facility with HCR contains operating and financial covenants that restrict our business and financing activities, and is subject to acceleration in specified circumstances, which may result in HCR taking possession and disposing of any collateral.** Under the terms of the RIFA, we may not, among other actions, without the prior written consent of HCR, (a) pay any dividends or make any other distribution or payment or redeem, retire or purchase any capital stock, except in certain prescribed circumstances, (b) create, incur, assume, or be liable with respect to any indebtedness except certain permitted indebtedness, or make or permit any payment on any indebtedness, except under certain limited circumstances, or (c) make any sale, transfer, out-license, lease or other disposition of any property or any economic interest, other than certain limited exceptions. Additionally, we are required (i) during the period from January 1, 2024 through December 31, 2024, to maintain at all times a minimum cash balance of \$7.5 million, and (ii) during all periods after December 31, 2024, to maintain at all times a minimum cash balance of \$15.0 million. Our obligations under the RIFA are collateralized by all of our assets and property, subject to limited exceptions. If we breach certain of our covenants in the RIFA and are unable to cure such breach within the prescribed period or are not granted waivers in relation to such breach, it may constitute an event of default under the RIFA, giving HCR the right to require us to repay the then outstanding obligations immediately, and HCR could, among other things, foreclose on the collateral granted to them to collateralize such indebtedness, which includes our intellectual property, if we are unable to pay the outstanding debt immediately. Our management has broad discretion in using the net proceeds from our financing facility with HCR and prior equity offerings and may not use them effectively. We are using the net proceeds of our financing facility with HCR, our September 2024 private equity offering, the September 2024 private placement, our December 2023 public equity offering, the December 2023 private placement and prior public and private equity offerings to support the development and commercialization of YUTREPIA, including the potential commercial launch of YUTREPIA in the event of final FDA approval, the commercialization of Treprostinil Injection, the development and servicing of pumps for the administration of Treprostinil Injection, the development of L606, and for general corporate purposes. Our management has broad discretion in the application of such proceeds and could spend the proceeds in ways that do not improve our results of operations or enhance the value of our equity. The failure by our management to apply these funds effectively could result in financial losses that could have a material adverse effect on our business, diminish cash flows available to service our obligations to HCR, cause the value of our equity to decline and delay the development of our product candidates. Pending their use, we may invest such proceeds in short-term, investment-grade, interest-bearing securities, which may not yield favorable returns. Our ability to use our net operating loss carryforwards and certain other tax attributes may be limited. Under Section 382 of the Internal Revenue Code of 1986, as amended (the "Code"), if a corporation undergoes an "ownership change", generally defined as a greater than 50.0% change (by value) in its equity ownership over a three-year period, the corporation's ability to use its pre-change net operating loss carryforwards and other pre-change tax attributes, such as research tax credits, to offset its post-change income may be limited. With our September 2024 public equity offering, the September 2024 private placement, the January 2024 private placement, our December 2023 public equity offering, the December 2023 private placement, our April 2022 public equity offering, our 2021 private placement, the closing of the RareGen acquisition in November 2020, our July 2020 public equity offering, our December 2019 private placement, issuances under our prior at-the-market facility, our March 2019 follow-on equity offering and our July 2018 initial public offering, as well as other past transactions, we may have already triggered an "ownership change" limitation. We have not completed a formal study to determine if any "ownership changes" within the meaning of IRC Section 382 have occurred. If "ownership changes" within the meaning of Section 382 of the Code have occurred, and if we earn net taxable income, our ability to use our net operating loss carryforwards and research and development tax credits generated since inception to offset U.S. federal taxable income may be subject to limitations, which could potentially result in increased future tax liability to us and could require us to pay U.S. federal income taxes earlier than would be required if such limitations were not in effect. Similar rules and limitations may apply for state income tax purposes. Changes to existing tax laws, or challenges to our tax positions could adversely affect our business and financial condition. The tax regimes to which we are subject or under which we operate are unsettled and may be subject to significant change. The issuance of additional guidance related to existing or future tax laws, or changes to tax laws or regulations proposed or implemented by the current or a future U.S. presidential administration, Congress, or taxing authorities in other jurisdictions could materially affect our tax obligations. For example, beginning in 2022, the Tax Cuts and Jobs Act of 2017 eliminated the option to deduct research and development expenditures in the year incurred and instead requires taxpayers to capitalize and subsequently amortize such expenditures over five years for research activities conducted in the United States and over 15 years for research activities conducted outside the United States. In January 2024, the U.S. House of Representatives passed the Tax Relief for American Families and Workers Act, which would retroactively

as to whether the bill will be enacted into law. As another example, in August 2022, the Inflation Reduction Act of 2022 was enacted, and, among other things, included a new 15% alternative minimum tax on the adjusted financial statement income of certain large corporations for tax years beginning after December 31, 2022. To the extent that such changes have a negative impact on us, including as a result of related uncertainty, these changes could adversely impact our business, results of operations and financial position. In addition, U.S. federal, state and local tax laws are extremely complex and subject to various interpretations. Although we believe that our tax estimates and positions are reasonable, there can be no assurance that our tax positions will not be challenged by relevant tax authorities. If the relevant tax authorities assess additional taxes on us, this could result in adjustments to, or impact the timing or amount of, taxable income, deductions or other tax allocations, which may adversely affect our results of operations and financial position.⁵⁰Table of ContentsWe are a late-stage clinical biopharmaceutical company with no approved products and no historical revenue from the sale of our own products, which may make it difficult for you to evaluate our business, financial condition and prospects. We are a late-stage clinical biopharmaceutical company with no history of commercial operations upon which you can evaluate our prospects other than the activities we have undertaken with respect to the Promotion Agreement with Sandoz. Drug product development involves a substantial degree of uncertainty. Our operations to date have been limited to engaging in promotional and non-promotional activities under the Promotion Agreement with Sandoz, developing our PRINT technology, undertaking preclinical studies and clinical trials for our product candidates and collaborating with pharmaceutical companies, including GSK, to expand the applications for our PRINT technology through licensing as well as joint product development arrangements. We have not obtained final marketing approval for any of our product candidates and, accordingly, have not demonstrated an ability to generate revenue from our own pharmaceutical products or successfully overcome the risks and uncertainties frequently encountered by companies undertaking drug product development. Consequently, your ability to assess our business, financial condition and prospects may be significantly limited. Further, the net losses that we incur may fluctuate significantly from quarter-to-quarter and year-to-year, such that a period-to-period comparison of our results of operations may not be a good indication of our future performance. Other unanticipated costs may also arise in connection with the development of our product candidates and commercialization of any approved products. A Liquidia PAH does not hold the FDA regulatory approval for Treprostinil Injection and is dependent on Sandoz to manufacture and supply Treprostinil Injection in compliance with FDA requirements, and is more broadly dependent on Sandoz's⁵¹ FDA and healthcare compliance relative to Treprostinil Injection. Sandoz holds the FDA approval, or the ANDA, for and controls Treprostinil Injection and is responsible among other things for the compliant manufacture, distribution, labeling, and advertising of Treprostinil Injection. As a result, we are dependent on Sandoz to manufacture and supply Treprostinil Injection, and are dependent on Sandoz for the continued FDA compliance of Treprostinil Injection. We do not have control over Sandoz's⁵¹ compliance with laws and regulations applicable to drug manufacturers and ANDA holders (for example, applicable current good manufacturing practices, or cGMPs; FDA labeling, promotional labeling, and advertising requirements; pharmacovigilance and adverse event reporting; and other ongoing FDA reporting and submission requirements), nor over its compliance with healthcare compliance and fraud, waste, and abuse laws, or similar regulatory requirements and other laws and regulations, such as those related to environmental health and safety matters. In addition, we have no control over the ability of Sandoz to maintain adequate quality control, quality assurance and qualified personnel, or other personnel with roles related to the regulatory compliance of Treprostinil Injection and its labeling, promotion, and advertising or of Sandoz's⁵¹ activities in relation to government healthcare programs. If the FDA or a comparable foreign regulatory authority finds deficiencies with the manufacture or quality assurance of Treprostinil Injection or identifies safety or efficacy concerns related to Treprostinil Injection, or if Sandoz otherwise is unable to comply with applicable laws, regulations and standards, Sandoz's⁵¹ ability to manufacture, sell and supply Treprostinil Injection could be limited.⁵² Sandoz's⁵¹ ability to consistently manufacture and supply Treprostinil Injection in a timely manner may also be interrupted by production shortages or other supply interruptions. Our share of net profits under the Promotion Agreement is reduced by certain manufacturing costs and other write-offs related to Sandoz's⁵¹ inability to sell Treprostinil Injection, including in the event that Treprostinil Injection expires prior to sale. Currently, Treprostinil Injection expires 24 months after the date of manufacture.⁵³ Sales of Treprostinil Injection are dependent on market acceptance of generic treprostinil for parenteral administration by patients, health care providers and by third-party payors, while interactions with these persons and entities are subject to compliance requirements. The commercial success of Treprostinil Injection may also be impacted by increasing generic competition which may result in declining prices for Treprostinil Injection. Our ability to sell Treprostinil Injection is dependent on market acceptance of generic treprostinil for parenteral administration by patients, health care providers and by third-party payors. If Treprostinil Injection does not achieve an adequate level of acceptance, we may not generate sufficient revenue to offset our cost of revenue.⁵⁴Table of ContentsAt the same time, arrangements with healthcare providers, physicians, third-party payors and customers, and our sales, marketing and educational activities, may expose us to broadly applicable fraud and abuse and other healthcare laws and regulations that may constrain our business or financial arrangements and relationships.⁵⁵ The degree of market acceptance of Treprostinil Injection will depend on a number of factors, including:⁵⁶ the efficacy, safety and potential advantages compared to alternative treatments;⁵⁷ our ability to offer Treprostinil Injection for sale at competitive prices (generic drug prices, after initial generic entry, have been observed to decline with the entrance of additional generic competition);⁵⁸ the convenience and ease of administration compared to alternative treatments;⁵⁹ product labeling or product insert requirements of the FDA or foreign regulatory authorities, including any limitations or warnings contained in a product's⁶⁰ approved labeling, including any black box warning;⁶¹ the willingness of the target patient population to try new treatments, including the generic version of a brand, and of physicians to prescribe such treatments;⁶² our ability to hire and retain sales and marketing personnel and their ability to support Sandoz under the Promotion Agreement;⁶³ the strength of Sandoz's⁶⁴ manufacturing and distribution support;⁶⁵ the requirement by third-party payors to use generic treprostinil for parenteral administration in place of Remodulin;⁶⁶ our ability to maintain availability of medical devices used to administer Treprostinil Injection and preferences of the target patient population and health care providers regarding the medical devices used to administer Treprostinil Injection versus medical devices used to administer Remodulin;⁶⁷ the availability of third-party coverage and adequate reimbursement for Treprostinil Injection;⁶⁸ the prevalence and severity of any side effects;⁶⁹ any restrictions on the use of Treprostinil Injection together with other medications;⁷⁰ our and Sandoz's⁷¹ ability to maintain relationships with the specialty pharmacies;⁷² and⁷³ the services provided by specialty pharmacies related to use of Treprostinil Injection.⁷⁴ Our business may also be impacted by the need to maintain compliant operations (including oversight and monitoring of personnel and our activities) in relation to interactions with the persons and parties noted above, relative to FDA and healthcare law requirements, and with consideration of government and industry compliance best practices. A Medical devices, which we do not control, are necessary for the administration of Treprostinil Injection. In order for Treprostinil Injection to be administered to patients, patients must use certain other medical equipment, including pumps, cartridges and infusion sets. We do not manufacture or control such medical equipment, which is manufactured by third parties and owned and dispensed by specialty pharmacies, hospitals or other third parties. Our ability to serve patients is dependent upon the ability of specialty pharmacies to maintain sufficient inventory of such medical equipment to provide to patients. If manufacturers cease to manufacture or support medical equipment or if specialty pharmacies are unable to obtain or maintain sufficient inventories of such medical equipment, our sales may be adversely impacted. We have worked with Chengdu to develop the RG Cartridge, which received FDA 510(k) clearance in March 2021. The ability of patients to administer Treprostinil Injection through subcutaneous injection is dependent on the continued availability of the RG Cartridge. If the RG Cartridge experiences any quality problems, recalls or other adverse events, our ability to provide Treprostinil Injection to patients who receive treprostinil through subcutaneous injection will be limited.⁷⁵Table of ContentsIn addition, to administer Treprostinil Injection through subcutaneous injection, patients currently must use the CADD-MS 3 infusion pump manufactured by ICU Medical. ICU Medical no longer manufactures the CADD-MS 3 infusion pump and has indicated that they will no longer support the CADD-MS 3 infusion pump. Although we believe that the number of available CADD-MS 3 infusion pumps will be sufficient to serve patients through 2025, it is possible that the availability of CADD-MS 3 infusion pumps could end earlier. Due to this limitation in the availability of pumps, specialty pharmacies will limit the number of patients that they place on subcutaneous Treprostinil Injection therapy in order to ensure that patients placed on subcutaneous administration of Treprostinil Injection will not have to discontinue such treatment due to the unavailability of pumps. Until we are able to obtain a pump to replace the CADD-MS 3, the number of patients that can receive subcutaneous administration of Treprostinil Injection will continue to be constrained, which would continue to adversely affect sales of Treprostinil Injection. We are seeking to work with third parties to develop or procure other pumps that can be used to administer Treprostinil Injection in the future. For example, we have entered into an agreement with Sandoz and Mainbridge to develop a new pump that can be used to administer Treprostinil Injection in the future. Such pumps will require FDA 510(k) clearance before they can be sold. There is no guarantee that we or our partners will receive FDA 510(k) clearance for any such pumps or, even if they do receive FDA 510(k) clearance for any such pumps, that they will do so in a timely manner. For example, we have still not submitted a 510(k) clearance application and are currently uncertain when, if ever, such a 510(k) clearance application will be submitted. If we are unable to identify, develop and obtain any required FDA clearance for new pumps for the subcutaneous administration of Treprostinil Injection prior to the unavailability of the CADD-MS 3, we may no longer be able to serve patients with Treprostinil Injection through the subcutaneous route of administration. Failure by us or third parties to successfully develop or supply the medical equipment or to obtain or maintain regulatory approval or clearance of such medical equipment could negatively impact the market acceptance of and sales of Treprostinil Injection. We maintain our cash at financial institutions, often in balances that exceed federally insured limits. Our cash is held in non-interest-bearing and interest-bearing accounts at multiple banking institutions that may exceed the Federal Deposit Insurance Corporation, or the FDIC, insurance limits. If such banking institutions were to fail, we could lose all or a portion of those amounts held in excess of such insurance limitations. For example, the FDIC took control of Silicon Valley Bank, where we previously held all of our cash and cash equivalents, on March 10, 2023. The Federal Reserve subsequently announced that account holders would be made whole, and we were able to move substantially all of our cash and cash equivalents to another financial institution. However, the FDIC may not make all account holders whole in the event of future bank failures. In addition, even if account holders are ultimately made whole with respect to a future bank failure, account holders' access to their accounts and assets held in their accounts may be substantially delayed. Any material loss that we may experience in the future or inability for a material time period to access our cash and cash equivalents could have an adverse effect on our ability to pay our operational expenses or make other payments, which could adversely affect our business. Risks Related to the Commercialization of our Product Candidates and Generic Treprostinil InjectionUnited Therapeutics has initiated lawsuits against us in which it claims that YUTREPIA is infringing its patents and that we have misappropriated its trade secrets and confidential information and has initiated a lawsuit against the FDA challenging the FDA's⁷⁶ acceptance of our amended NDA for YUTREPIA for review, which may result in our company being further delayed in its efforts to commercialize YUTREPIA and may limit the indications for which YUTREPIA is approved. We are developing YUTREPIA under the 505(b)(2) regulatory pathway with Tyvaso as the reference listed drug. Accordingly, under the Hatch-Waxman Amendments to the Food, Drug and Cosmetic Act, we were required to, in the NDA for YUTREPIA, certify that patents listed in the Orange Book for Tyvaso are invalid, unenforceable or will not be infringed by the manufacture, use or sale of YUTREPIA.⁷⁷Table of ContentsIn connection with an amendment to our NDA filed in July 2023 to add PH-ILD as an indication for YUTREPIA, we provided a new notice of the paragraph IV certification to United Therapeutics as the owner of the patents that are the subject of the certification to which the NDA for YUTREPIA refers. As a result, in September 2023, United Therapeutics filed a complaint for patent infringement against us in the U.S. District Court for the District of Delaware (Case No. 1:23-cv-00975-RGA) (the "New Hatch-Waxman Litigation"). In the New Hatch-Waxman Litigation, United Therapeutics is asserting that the Company infringes U.S. Patent No. 11,826,327, or the "327 Patent", entitled "Treatment for Interstitial Lung Disease".⁷⁸ In February 2024, United Therapeutics filed a motion seeking a preliminary injunction to prevent us from manufacturing, marketing, storing, importing, distributing, offering for sale, and/or selling YUTREPIA for the treatment of PH-ILD. Judge Andrews denied the motion for a preliminary injunction in May 2024. Discovery in the case remains ongoing. Although we do not believe United Therapeutics is entitled to a new 30-month stay or a preliminary injunction in connection with the New Hatch-Waxman Litigation, it is possible that the Court could rule that a new mandatory 30-month delay has been triggered with respect to the approval of the 505(b)(2) NDA application or that a preliminary injunction is warranted. In February 2024, United Therapeutics also filed a lawsuit against the FDA, challenging the FDA's⁷⁹ acceptance of our amended NDA for review (the "Original FDA Litigation"). In March 2024, United Therapeutics filed a motion for a temporary restraining order in the Original FDA Litigation, seeking to enjoin the FDA from approving our NDA for YUTREPIA with respect to the indication to treat PH-ILD. United Therapeutics' motion was denied in March 2024. In May 2024, both we and the FDA filed motions to dismiss United Therapeutics'⁸⁰ complaint. Prior to the Court's⁸¹ ruling on the motions to dismiss, United Therapeutics voluntarily dismissed its complaint in the Original FDA Litigation without prejudice. In September 2024, United Therapeutics re-asserted its challenge to FDA's⁸² acceptance of our amended NDA for review as a cross claim in the lawsuit we instituted against the FDA in August 2024 (the "New FDA Litigation"). Although we do not believe the arguments of United Therapeutics have merit, it is possible that the Court could rule that the FDA must reject the amendment to the YUTREPIA NDA to add PH-ILD to the label, in which case we may be required to later file a supplement to our NDA to add PH-ILD to the label. If we are required to file a supplement to add PH-ILD to the label for YUTREPIA, although we do not believe United Therapeutics would be entitled to a new 30-month stay, it is possible that the FDA or a Court could rule that a new mandatory 30-month delay has been triggered with respect to the supplement. In addition, United Therapeutics may seek to assert newly issued patents against us, including U.S. Patent Number 11,723,887, and may seek to enjoin the FDA from granting final approval to YUTREPIA or enjoin us from launching YUTREPIA through one or more additional legal proceedings. As a result of this litigation instituted to date and potential litigation that may be instituted in the future, we may be subject to significant delay and incur substantial additional costs in litigation before we are able to commercialize YUTREPIA, if at all. In addition, if United Therapeutics is successful in any of its claims that it has brought to date or any claims it may bring in the future, we may be unable to commercialize YUTREPIA for the treatment of one or more indications or at all until the expiration of the applicable United Therapeutics patents, which could materially harm our business. For example, in the event United Therapeutics prevails with respect to its claims regarding the "327 Patent", it is possible that an injunction could be issued, preventing the FDA from granting final approval for YUTREPIA for PH-ILD or forcing the FDA to revoke any prior approval for YUTREPIA for PH-ILD. Also, although United Therapeutics' initial requests for injunctive relief have been denied, if United Therapeutics is successful in obtaining a preliminary injunction or temporary restraining order in the New Hatch-Waxman Litigation or the New FDA Litigation, we could be limited to commercializing YUTREPIA only for the PAH indication for an extended time period. In December 2021, United Therapeutics filed a complaint in the Superior Court in Durham County, North Carolina, alleging that we and Robert Roscigno (the "Roscigno"), a former United Therapeutics employee, who later joined us as an employee many years after terminating his employment with United Therapeutics, conspired to misappropriate certain trade secrets of United Therapeutics and engaged in unfair or deceptive trade practices. In January 2024, Dr. Roscigno filed a motion for summary judgment on all claims, but the motion was denied in July 2024. In addition, in July 2024, the Company filed a motion for summary judgment with respect to all claims. Briefing on the Company's⁸³ motion is complete and a hearing has been scheduled for December 19, 2024.⁸⁴Table of ContentsIn May 2024, United Therapeutics filed a second complaint in the Superior Court in Durham County, North Carolina, against Dr. Roscigno, alleging that he breached prior employment agreements with United Therapeutics by failing to assign to United Therapeutics his interest in patents obtained by the Company that relied upon or benefited from certain inventions, discoveries, materials, authorship, derivatives and results developed by Dr. Roscigno while he was employed by United Therapeutics. The Company was also named as a defendant in this new lawsuit. As part of the lawsuit, United Therapeutics alleges that Dr. Roscigno misappropriated certain intellectual property of United Therapeutics which led to the development of YUTREPIA. The complaint also seeks declaratory judgement such that all right, title and interest in and to any patentable or unpatentable inventions, discoveries, and ideas made or conceived by Dr. Roscigno while employed by the Company should be assigned and transferred to United Therapeutics because they involved the use of United Therapeutics' confidential information. On July 30, 2024, the Company filed a motion to dismiss all claims. Briefing on the motion is complete and a hearing has been scheduled for December 19, 2024. Success in a lawsuit, including in any such lawsuit with respect to some patents or some claims in a given patent, does not mean that we will be similarly successful upon appeal of those decisions. In addition, success in one proceeding, including with respect to a given patent, patent claim or trade secret in one proceeding, does not mean we will be similarly successful with respect to that same or a similar patent, patent claim or trade secret in another proceeding. If we are found to infringe, misappropriate or otherwise violate any United Therapeutics' intellectual property rights, we could be required to obtain a license from United Therapeutics to continue developing and marketing YUTREPIA. However, we may not be able to obtain any required license on commercially reasonable terms or at all. We could be found liable for monetary damages, including treble damages and attorneys' fees if we are found to have willfully infringed a patent or to have misappropriated a trade secret of United Therapeutics. In addition, we may be forced to

redesign YUTREPIA to avoid infringement. We face significant competition from large pharmaceutical companies, among others, in developing our products and in gaining regulatory approval to bring them to market in time to achieve commercial success, and our operating results will suffer if we are unable to compete effectively. We face significant competition from industry players worldwide, including large multi-national pharmaceutical companies, other emerging or smaller pharmaceutical companies, as well as universities and other research institutions. Many of our competitors have substantially greater financial, technical and other resources, such as a larger research and development staff and more experience in manufacturing and marketing, than we do. As a result, these companies may obtain marketing approval for their product candidates more quickly than we are able to and/or be more successful in commercializing their products, including generic treprostinal products, than us. Smaller or early-stage companies may also prove to be significant competitors, particularly through collaboration arrangements with large, established companies. We may also face competition as a result of advances in the commercial applicability of new technologies and greater availability of capital for investment in such technologies. Our competitors may also invest heavily in the discovery and development of novel drug products that could make our product candidates less competitive or may file FDA citizen petitions or other correspondence with the FDA, as United Therapeutics has recently done, which may delay the approval process for our product candidates. Furthermore, our competitors may succeed in developing, acquiring or licensing, on an exclusive basis, pharmaceutical products that are easier to develop, more effective or less costly than any product candidates that we are currently developing or that we may develop. Our competitors may also succeed in asserting existing patents or developing new patents, including patents that may issue from patent applications that are currently being pursued by United Therapeutics, to which we do not have a license, in an attempt to prevent us from marketing our products. These competitors may also compete with us in recruiting and retaining qualified sales personnel. Any new drug product that competes with a prior approved drug product must demonstrate advantages in safety, efficacy, tolerability or convenience in order to overcome price competition and to be commercially successful. Our products, if and when approved, are expected to face competition from drug products that are already on the market, as well as those in our competitors' development pipelines. We expect that our lead program, YUTREPIA, an inhaled treprostinal therapy for the treatment of PAH and PH-ILD, and L606, a nebulized, liposomal formulation of treprostinal for treatment of PAH and PH-ILD, will face competition from the following inhaled prostacyclin analog therapies that are either currently marketed or in clinical development: 55Table of Contents—Tyvaso (treprostinal), marketed by United Therapeutics, has been approved for the treatment of PAH in the United States since 2009 and for PH-ILD since 2021. Tyvaso is the reference listed drug in our NDA for YUTREPIA. Following patent litigation, United Therapeutics and Watson Pharmaceuticals reached a settlement whereby Watson Pharmaceuticals will be permitted to enter the market with a generic version of Tyvaso beginning on January 1, 2026. Tyvaso DPI (treprostinal), licensed from MannKind by United Therapeutics, is a dry-powder formulation of treprostinal that was approved for the treatment of PAH and PH-ILD in the United States in May 2022. TIPiP (Tremipalmitin Inhalation Powder (TPIP)), a dry-powder formulation of a treprostinal prodrug being developed by Insmed. Insmed announced the completion of an initial Phase 1 study in February 2021 which demonstrated that TPIP was generally safe and well tolerated, with a pharmacokinetic profile that supports once-daily dosing. Insmed initiated Phase 2 trials studying patients diagnosed with PAH and PH-ILD in May 2021 and December 2022, respectively. In May 2024, Insmed reported positive topline safety and tolerability data as well as certain exploratory efficacy endpoints from the Phase 2 PH-ILD. Based on these Phase 2 results, Insmed is pursuing discussions with global regulatory authorities on the design of a Phase 3 study in PH-ILD to initiate in 2025. If the TPIP clinical program is successful in demonstrating less frequent dosing with similar efficacy and safety to YUTREPIA and Tyvaso DPI, then TPIP has the potential to be viewed as a more attractive option and may take market share rapidly. Ventavis® (iloprost), marketed by Actelion, a division of Johnson & Johnson, has been approved for the treatment of PAH in the United States since 2004. In addition to these other inhaled treprostinal therapies, we expect that YUTREPIA and L606 will also face competition from other treprostinal-based drugs, including Orenitram, which is administered orally, and Remodulin, which is administered parenterally, both of which are marketed by United Therapeutics. Branded pharmaceutical companies such as United Therapeutics continue to defend their products vigorously through, among other actions, life cycle management, marketing agreements with third-party payors, pharmacy benefit managers and generic manufacturers. These actions add increased competition in the generic pharmaceutical industry, including competition for Treprostinal Injection. Additionally, even though Sandoz launched the first-to-file fully substitutable generic treprostinal for parenteral administration in March 2019 that is sold primarily through the specialty pharmacies, Teva Pharmaceutical Industries Ltd. launched a generic treprostinal for parenteral administration in October 2019 that is sold primarily through a specialty pharmacy and to hospitals, Par Pharmaceutical, Inc. launched a generic treprostinal for parenteral administration after receiving approval in September 2019 that is sold primarily to hospitals, Dr. Reddy's Laboratories Inc. launched a generic treprostinal for parenteral administration in April 2023, and Alembic received approval in February 2021 for generic treprostinal for parenteral administration. Such increased competition may result in a smaller than expected commercial opportunity for us. Generic drug prices may, and often do, decline, sometimes dramatically, especially as additional generic pharmaceutical companies (including low-cost generic producers outside of the United States) receive approvals and enter the market for a given product. The goals established under the Generic Drug User Fee Act, and increased funding of the FDA's Office of Generic Drugs, have led to more and faster generic approvals, and consequently increased competition for generic products. The FDA has stated that it has established new steps to enhance competition, promote access and lower drug prices and is approving record-breaking numbers of generic applications. The FDA's changes may benefit our competitors. Our ability to sell Treprostinal Injection and earn revenue is affected by the number of companies selling competitive products, including new market entrants, and the timing of their approvals. 56Table of Contents In addition to treprostinal-based therapies, other classes of therapeutic agents for the treatment of PAH include the following: IP-agonists, such as sifelipag, marketed by Actelion, and ralinepag, licensed from Arena Pharmaceuticals, Inc. by United Therapeutics, which is currently in clinical development. Endothelin receptor antagonists, such as bosentan and macitentan, both marketed by Actelion, and ambrisentan, marketed by Gilead. Generic versions of bosentan and ambrisentan are currently available. PDE-5 inhibitors, such as tadalafil, marketed by United Therapeutics, and sildenafil, marketed by Pfizer Inc. Generic versions of both tadalafil and sildenafil are currently available. Soluble guanylate cyclase (sGC) stimulator, such as oral riociguat marketed by Bayer for PAH, and inhaled mosciguat being developed by Pulmovant for PH-ILD. Activin signaling inhibitor, such as sotatercept marketed by Merck & Co. and KER-012 being developed by Keros Therapeutics/Merck & Co. injectable sotatercept, with a brand name of Winrevair, was approved by the FDA in March 2024 and is a potential first-in-class molecule that targets the proliferation of cells in the pulmonary arterial wall. Its clinical use is developing, and it is possible that it may be used prior to prostacyclin therapies, which may have an adverse effect on the market potential for YUTREPIA and/or L606. We are also aware of several other agents in clinical development that are exploring mechanisms of action which, if approved, could impact the standard of care for treating PAH and/or PH-ILD in the United States, including programs from Merck & Co. Inc., and Gossamer Bio, Inc., among others. There are a number of competitors seeking marketing approval and/or regulatory exclusivity with respect to products that are or would be competitive to our product candidate. Thus, we face the risk that one of our competitors will be granted marketing approval and/or regulatory exclusivity before we are able to obtain FDA approval for our product candidate. In that case, as stated above, there is the possibility that such a competitor would be able to prevent us from obtaining approval of and marketing our product candidate until the expiration of the competitor's term of FDA regulatory exclusivity, which could be a term of three years for so-called New Clinical Investigation exclusivity, or could conceivably be for longer periods of time if the competitor is successful in being granted other forms of FDA regulatory exclusivity which might include, for example, Orphan Disease Designation exclusivity (seven years), New Chemical Entity exclusivity (five years), or Pediatric exclusivity (six months beyond other existing exclusivities or patent terms). For example, United Therapeutics was recently awarded New Clinical Investigation exclusivity for Tyvaso DPI, which will expire in May 2025. As a result, unless we are successful in having such exclusivity overturned, the FDA will be unable to approve YUTREPIA until after the exclusivity expires in May 2025. In the event United Therapeutics sought and was able to obtain one or more other regulatory exclusivities with respect to Tyvaso DPI, it could further significantly delay our ability to obtain final approval for YUTREPIA. Even if the FDA does not recognize any new regulatory exclusivity for United Therapeutics, United Therapeutics could challenge the FDA's decision and seek an injunction to prevent approval of YUTREPIA in one or more indications until such challenge has been decided. In addition, if one of our competitors is granted marketing approval before we are able to obtain FDA approval for our product candidates, as was the case with respect to the approval of United Therapeutics' Tyvaso DPI product, such competitors will be able to promote and market their products before we are able to do so, which may place us at a competitive disadvantage in the marketplace. One or more products that are competitive with YUTREPIA could also obtain approval for additional indications or broader conditions of use. These additional indications and broader conditions of use could be protected by one or more patents or regulatory exclusivities, preventing YUTREPIA from obtaining approval for the same indications or conditions of use. For instance, if Liquida is prevented from launching or selling YUTREPIA for the treatment of PH-ILD in connection with the patent litigation related to the '327 patent or the lawsuit that United Therapeutics filed 57Table of Contents against the FDA, Tyvaso and Tyvaso DPI would have broader labels than YUTREPIA. In addition, United Therapeutics is currently studying Tyvaso for the treatment of idiopathic pulmonary fibrosis, an indication for which it has received an orphan drug designation. Thus, even if YUTREPIA is approved, such competitive products could have a broader label than the initial label for YUTREPIA. If YUTREPIA has a narrower label than other competitive products, it may affect our ability to compete with such products. The ability of competitors to utilize other regulatory incentive programs could also expedite their FDA review and approval timeline, which could result in their products reaching the market before our product candidate, and which could create further potential implications on exclusivity as noted above. For example, when a Priority Review Voucher is redeemed in connection with an NDA, the FDA's goal review period would generally be expedited to six months, although this timeframe is not guaranteed. If we are unable to maintain our competitive position, our business and prospects will be materially and adversely affected. Our products may not achieve market acceptance or adequate third-party payor coverage. We are currently focused on developing drug products that can be approved under abbreviated regulatory pathways in the United States, such as the 505(b)(2) regulatory pathway, which allows us to rely on existing knowledge of the safety and efficacy of the relevant reference listed drugs to support our applications for approval in the United States. While we believe that it will be less difficult for us to convince physicians, patients and other members of the medical community to accept and use our drug products as compared to entirely new drugs, our drug products may nonetheless fail to gain sufficient market acceptance by physicians, patients, other healthcare providers and third-party payors. If any of our drug products fail to achieve sufficient market acceptance or third-party payor coverage, we may not be able to generate sufficient revenue to become profitable. The degree of market acceptance and third-party payor coverage of our drug products, if and when they are approved for commercial sale, will depend on a number of factors, including but not limited to: the timing of our receipt of marketing approvals, the terms of such approvals and the countries in which such approvals are obtained; the safety, efficacy, reliability and ease of administration of our drug products; the prevalence and severity of undesirable side effects and adverse events; the extent of the limitations or warnings required by the FDA or comparable regulatory authorities in other countries to be contained in the labeling of our drug products; the clinical indications for which our drug products are approved; the availability and perceived advantages of alternative therapies; any publicity related to our drug products or those of our competitors; the quality and price of competing drug products; our ability to obtain third-party payor coverage and sufficient reimbursement; the willingness of patients to pay out of pocket in the absence of third-party payor coverage; and the selling efforts and commitment of our commercialization collaborators. If our drug products, if and when approved, fail to receive a sufficient level of market acceptance or sufficient third-party payor coverage, our ability to generate revenue from sales of our drug products will be limited, and our business and results of operations may be materially and adversely affected. We may not be able to build a commercial operation, including establishing and maintaining marketing and sales capabilities or entering into agreements with third parties to market and sell our drug products. In order to market and sell any of our drug products, if and when approved, we will be required to build our marketing and sales capabilities with respect to such products. With the acquisition of Liquida PAH, we acquired a sales force to market generic treprostinal in accordance with the Promotion Agreement. In addition, during 2023, we significantly increased the size of our sales force in anticipation of a potential launch of YUTREPIA. However, if we experience 58Table of Contents continued delays in the approval of YUTREPIA, we may be unable to retain our sales force. Moreover, we cannot assure you that we will be successful in further building or effectively managing our marketing and sales capabilities or be able to do so in a cost-effective manner. In addition, we may enter into collaboration arrangements with third parties to market our drug products. We may face significant competition for collaborators. In addition, collaboration arrangements may be time-consuming to negotiate and document. We cannot assure you that we will be able to negotiate collaborations for the marketing and sales of our drug products on acceptable terms, or at all. Even if we do enter into such collaborations, we cannot assure you that our collaborators will be successful in commercializing our products. If we or our collaborators are unable to successfully commercialize our drug products, whether in the United States or elsewhere, our business and results of operations may be materially and adversely affected. As we seek to establish a commercial operation with respect to YUTREPIA in anticipation of potential approval from the FDA, we also continue to evaluate and develop additional drug candidates, including L606. There can be no assurance that we will be able to successfully manage the balance of our research and development operations with our commercial activities. Potential investors should be aware of the problems, delays, expenses and difficulties frequently encountered by companies balancing development of product candidates, which can include problems such as unanticipated issues relating to clinical trials and receipt of approvals from the FDA and foreign regulatory bodies, with commercialization efforts, which include problems relating to managing manufacturing and supply, reimbursement, marketing problems, and other additional costs. There are risks involved with building and expanding our sales, marketing, and other commercialization capabilities. For example, recruiting and training a sales force is expensive and time-consuming. If the commercial launch of a drug candidate for which we recruit or have recruited a sales force and establish marketing capabilities is delayed or does not occur for any reason, we would have prematurely or unnecessarily incurred these commercialization expenses. This may be costly, and our investment would be lost if we cannot retain or reposition our sales and marketing personnel. Factors that may impact our efforts to commercialize our drug candidates on our own and generate product revenues include: our inability to recruit and retain adequate numbers of effective sales and marketing personnel over a large geographic area; the costs and time associated with the initial and ongoing training of sales and marketing personnel on legal and regulatory compliance matters and monitoring their actions; understanding and training relevant personnel on the limitations on, and the transparency and reporting requirements applicable to, remuneration provided to actual and potential referral sources; the clinical indications for which the products are approved and the claims that we may make for the products; limitations or warnings, including distribution or use restrictions, contained in the products' approved labeling; the inability of sales personnel to obtain access to physicians or to effectively promote any future drugs; the lack of complementary drugs to be offered by sales personnel, which may put us at a competitive disadvantage relative to companies with more extensive product lines; any distribution and use restrictions imposed by the FDA or to which we agree; liability for sales and marketing personnel who fail to comply with the applicable legal and regulatory requirements; our ability to maintain a healthcare compliance program including effective mechanisms for compliance monitoring; and unforeseen costs and expenses associated with creating a sales and marketing organization. In the future, we may choose to participate in sales activities with collaborators for some of our drug candidates. However, there are also risks with entering into these types of arrangements with third parties to perform sales, marketing and distribution services. For example, we may not be able to enter into such arrangements on terms that are favorable to us. Our drug revenues or the profitability of these drug revenues to us are likely to be lower than if we were to market and sell any drug candidates that we develop ourselves. In addition, we likely will have little control over such third parties, and any of them may fail to devote the necessary resources and attention to sell and market our drug 59Table of Contents candidates effectively. If we do not establish sales and marketing capabilities successfully, either on our own or in collaboration with third parties, we will not be successful in commercializing our drug candidates. Further, our business, results of operations, financial condition and prospects will be materially adversely affected. We may be exposed to claims and may not be able to obtain or maintain adequate product liability insurance. Our business is exposed to the risk of product liability and other liability risks that are inherent in the development, manufacture, clinical testing, commercialization and marketing of pharmaceutical products. These risks exist even if a product is approved for commercial sale by the FDA or comparable regulatory authorities in other countries and manufactured in licensed facilities. Our current product candidates, YUTREPIA and L606, and Treprostinal Injection are designed to affect important bodily functions and processes. Any side effects, manufacturing defects, misuse or abuse associated with our products could result in injury to a patient or even death. Claims that are successfully brought against us could have a material and adverse effect on our financial condition and results of operations. Further, even if we are successful in defending claims brought against us, our reputation could suffer. Regardless of merit or eventual outcome, product liability claims may also result in, among others: a decreased demand for our products; a withdrawal or recall of our products from the

market; a withdrawal of participants from our ongoing clinical trials; the distraction of our management's attention from our core business activities to defend such claims; additional costs to us; and a loss of revenue. Our insurance may not provide adequate coverage against our potential liabilities. Furthermore, we, our collaborators or our licensees may not be able to obtain or maintain insurance on acceptable terms, or at all. Our inability to obtain sufficient product liability insurance at an acceptable cost and/or scope of coverage to protect against potential product liability claims could prevent or inhibit the commercialization of products we develop, alone or with our collaborators. The market for insurance coverage is increasingly expensive, and the costs of insurance coverage will increase as our clinical programs and commercialization efforts increase in size. In addition, our collaborators or licensees may not be willing to indemnify us against these types of liabilities and may not themselves be sufficiently insured or have sufficient assets to satisfy any product liability claims. To the extent that they are uninsured or uninsurable, claims or losses that may be suffered by us, our collaborators or our licensees may have a material and adverse effect on our financial condition and results of operations. Any claims against us, regardless of their merit, could severely harm our financial condition, strain our management and other resources, adversely affect or eliminate the prospects for commercialization or sales of a product that is the subject of any such claim, and could have a material adverse effect on our business, financial condition, results of operations, and growth prospects. Risks Related to the Development and Regulatory Approval of our Product Candidates We are primarily dependent on the success of our product candidate, YUTREPIA, for which we received tentative approval from the FDA, and this product candidate may fail to receive final marketing approval (in a timely manner or at all), may fail to receive approval for one or more indications for which we have sought approval or may not be commercialized successfully. We do not have any products approved for marketing in any jurisdiction and we have never generated any revenue from sales of our own products. Our ability to generate revenue from sales of our own products and achieve profitability depends on our ability, alone or with strategic collaboration partners, to successfully complete the development of, and obtain the regulatory and marketing approvals necessary to commercialize, one or more of our product candidates. We expect that a substantial portion of our efforts and expenditure over the next few years will be devoted to our product candidate, YUTREPIA, a proprietary inhaled dry powder formulation of treprostinil for the treatment of PAH and PH-ILD, and L606, a nebulized, liposomal formulation of treprostinil for treatment of PAH and PH-ILD. 60Table of Contents We received tentative approval of our NDA for YUTREPIA for the treatment of PAH and PH-ILD in August 2024. However, our receipt of tentative approval does not mean that we will receive final approval of our NDA for YUTREPIA in a timely manner or at all or that we will receive final approval for both indications. United Therapeutics has invested considerable time and resources in an effort to block final approval of YUTREPIA, and expectations related to final FDA approval and projected product launch timelines are impacted by ongoing litigation following lawsuits filed by United Therapeutics. For instance, in connection with an amendment to our NDA filed on July 24, 2023 to add PH-ILD as an indication for YUTREPIA, we provided a new notice of the paragraph IV certification to United Therapeutics as the owner of the patents that are the subject of the certification to which the NDA for YUTREPIA refers. As a result, in September 2023, United Therapeutics filed the New Hatch-Waxman Litigation, again asserting infringement by the Company of the '793 Patent, which lawsuit was amended on November 30, 2023, to add claims asserting infringement of the '327 Patent. Although the claims related to the '793 Patent were subsequently withdrawn, in February 2024, United Therapeutics also filed a motion seeking a preliminary injunction to prevent us from manufacturing, marketing, storing, importing, distributing, offering for sale, and/or selling YUTREPIA for the treatment of PH-ILD. That motion for preliminary injunction was denied, but United Therapeutics may still seek injunctive relief in the future. In September 2024, United Therapeutics also filed a cross claim in the New FDA Litigation, seeking to enjoin the FDA from approving our NDA for YUTREPIA with respect to the indication to treat PH-ILD. Although we do not believe United Therapeutics is entitled to any injunction or temporary restraining order in the New Hatch-Waxman Litigation or the New FDA Litigation, it is possible that the Court could rule that the FDA must reject the amendment to the YUTREPIA NDA to add PH-ILD to the label or that, even if YUTREPIA has launched for both PAH and PH-ILD, the Company must remove PH-ILD from the label for YUTREPIA. In addition, a drug product that is granted tentative approval, like YUTREPIA, may be subject to additional review before final approval, particularly if tentative approval was granted more than three years before the earliest lawful approval date. The FDA's tentative approval of YUTREPIA for the treatment of PAH and PH-ILD was based on information available to FDA at the time of the tentative approval letter (i.e., information in the application and the status of current good manufacturing practices of the facilities used in the manufacturing and testing of the drug product) and is therefore subject to change on the basis of new information that may come to FDA's attention. A new drug product may not be marketed until the date of final approval. Expectations for YUTREPIA and/or L606 also may be impacted by competing products, including Tyvaso® DPI. See Item 1A. Risk Factors. "We face significant competition from large pharmaceutical companies, among others, in developing our products and in gaining regulatory approval to bring them to market in time to achieve commercial success, and our operating results will suffer if we are unable to compete effectively. We cannot assure you that we will receive final marketing approval for YUTREPIA or L606 or, even if we do receive final marketing approval, the indications for which they will be approved. The FDA or comparable regulatory authorities in other countries may delay, limit or deny final approval of our product candidate for various reasons. For example, such authorities may disagree with the design, scope or implementation of our clinical trials, or with our interpretation of data from our preclinical studies or clinical trials. Further, there are numerous FDA personnel assigned to review different aspects of an NDA, and uncertainties can be presented by their ability to exercise judgment and discretion during the review process. During the course of review prior to final approval, the FDA may request or require additional preclinical, clinical, chemistry, manufacturing, and control (CMC) or other data and information or conduct additional inspections. If any additional issues were identified in such information requests or inspections or if FDA determines that we failed to include required CMC information in the NDA for our products, including YUTREPIA, we may be delayed in obtaining final approval or may be unable to obtain final approval. Furthermore, responses to FDA's requests may be time-consuming and expensive. Status as a combination product, as is the case for YUTREPIA and L606, may complicate or delay the FDA review process. Product candidates that the FDA deems to be combination products, such as YUTREPIA and L606, or that otherwise rely on innovative drug delivery systems, may face additional challenges, risks and delays in the product development and regulatory approval process. Additionally, the FDA could delay approval of YUTREPIA and/or L606 even if approved after completing its review. For example, Tyvaso DPI was granted regulatory exclusivity that will delay final approval of YUTREPIA until after the exclusivity expires in May 2025. If a competing product comprised of an inhaled dry-powder formulation of treprostinil, such as Tyvaso DPI, is granted additional regulatory exclusivity, that could delay the final approval of YUTREPIA until said exclusivity expires. Moreover, the applicable requirements for approval may differ from country to country. It is also possible that 61Table of Contents recent decisions by the United States Supreme Court, eliminating court deference to decisions by administrative agencies, may delay any final decisions from the FDA as it considers how to implement this new ruling into its decision-making process. If we successfully obtain marketing approvals for YUTREPIA and/or L606, we cannot assure you that they will be commercialized in a timely manner or successfully, or at all. For example, even if such products are approved by the FDA, they may not achieve a sufficient level of market acceptance or third-party payor coverage, or we may not be able to effectively build our marketing and sales capabilities or scale our manufacturing operations to meet commercial demand. The successful commercialization of YUTREPIA and L606 will also, in part, depend on factors that are beyond our control. Therefore, we may not generate significant revenue from the sale of such products, even if approved. Any delay or setback we face in the commercialization of YUTREPIA and/or L606 may have a material and adverse effect on our business and prospects, which will adversely affect your investment in our company. Our preclinical studies and clinical trials may not be successful and delays in such preclinical studies or clinical trials may cause our costs to increase and significantly impair our ability to commercialize our product candidates. Results of previous clinical trials or interim results of ongoing clinical trials may not be predictive of future results. Before we are able to commercialize our drug products, we are required to undertake extensive preclinical studies and clinical trials to demonstrate that our drug products are safe and effective for their intended uses. However, we cannot assure you that our drug products will, in preclinical studies and clinical trials, demonstrate safety and efficacy as necessary to obtain marketing approval. Due to the nature of drug product development, many product candidates, especially those in early stages of development, may be terminated during development. Although we believe we have completed clinical development for YUTREPIA, we have not yet obtained final approval for or commercialized any of our own product candidates and as a result do not have a track record of successfully bringing our own product candidates to market. Furthermore, YUTREPIA and L606 have, to date, been tested only in relatively small study populations and, accordingly, the results from our earlier clinical trials may be less reliable than results achieved in larger clinical trials, if required. Additionally, the outcome of preclinical testing and early clinical trials may not be predictive of the success of later clinical trials, and preliminary and interim results of a clinical trial do not necessarily predict final results. Preclinical studies and clinical trials may fail due to factors such as flaws in trial design, dose selection and patient enrollment criteria. The results of preclinical studies and early clinical trials may not be indicative of the results of subsequent clinical trials. Product candidates may, in later stages of clinical testing, fail to show the desired safety and efficacy traits despite having progressed through preclinical studies and earlier clinical trials. Moreover, there may be significant variability in safety or efficacy results between different trials of the same product candidate due to factors including, but not limited to, changes in trial protocols, differences in the composition of the patient population, adherence to the dosing regimen and other trial protocols and amendments to protocols and the rate of drop-out among patients in a clinical trial. If our preclinical studies or clinical trials are not successful and we are unable to bring our product candidates to market as a result, our business and prospects may be materially and adversely affected. Furthermore, conducting preclinical studies and clinical trials is a costly and time-consuming process. The length of time required to conduct the required studies and trials may vary substantially according to the type, complexity, novelty and intended use of the product candidate. A single clinical trial may take up to several years to complete. Moreover, our preclinical studies and clinical trials may be delayed or halted due to various factors, including, among others: delays in raising the funding necessary to initiate or continue a clinical trial; delays in manufacturing sufficient quantities of product candidates for clinical trials; delays in reaching agreement on acceptable terms with prospective contract research organizations (CROs) and clinical trial sites; delays in obtaining institutional review board approval at clinical trial sites; delays in recruiting suitable patients to participate in a clinical trial; delays in patients. A completion of clinical trials or their post-treatment follow-up; regulatory authorities' interpretation of our preclinical and clinical data; and 62Table of Contents unforeseen safety issues, including a high and unacceptable severity, or prevalence, of undesirable side effects or adverse events caused by our product candidates or similar drug products or product candidates. If our preclinical studies or clinical trials are delayed, the commercialization of our product candidates will be delayed and, as a result, we may incur substantial additional costs or not be able to recoup our investment in the development of our product candidates, which would have a material and adverse effect on our business. Clinical trials and data analysis can be expensive, time-consuming and difficult to design and implement. If we are unsuccessful in obtaining regulatory approval for our products, or any required clinical studies of our products do not provide positive results, we may be required to delay or abandon development of such products, which would have a material adverse impact on our business. Continuing product development requires additional and extensive clinical testing. Human clinical trials are very expensive and difficult to design and implement, in part because they are subject to rigorous regulatory requirements. The clinical trial process is also time-consuming. We cannot provide any assurance or certainty regarding when we might receive regulatory approval for our products, including YUTREPIA and L606. Furthermore, failure can occur at any stage of the process, and we could encounter problems that cause us to abandon an NDA filed with the FDA or repeat clinical trials. The commencement and completion of clinical trials for any current or future development product candidate may be delayed by several factors, including: unforeseen safety issues; determination of dosing issues; lack of effectiveness during clinical trials; slower than expected rates of patient recruitment; inability to monitor patients adequately during or after treatment; and inability or unwillingness of medical investigators to follow our clinical protocols or amendments to our protocols. In addition, the FDA or an independent IRB may suspend our clinical trials at any time if it appears that we are exposing participants to unacceptable health risks or if the FDA finds deficiencies in our IND submissions or the conduct of these trials. Therefore, we cannot provide any assurance or predict with certainty the schedule for future clinical trials. Although clinical data is an essential part of NDA filings, NDAs must also contain a range of additional data including CMC data to meet FDA standards for approval. In the event we do not ultimately receive final regulatory approval for YUTREPIA and/or L606, we may be required to terminate development of these product candidates. The marketing approval processes of the FDA and comparable regulatory authorities in other countries are unpredictable and our product candidates may be subject to multiple rounds of review or may not receive marketing approval. Pursuing marketing approval for a pharmaceutical product candidate (for example, through the NDA process) is an extensive, lengthy, expensive and inherently uncertain process. We cannot assure you that any of our product candidates will receive marketing approval. Regulatory authorities may delay, limit or deny approval of our product candidates for many reasons, including, but not limited to, the following: the FDA or comparable regulatory authorities may, for a variety of reasons, take the view that the data collected from our preclinical and clinical trials and human factors testing, or data that we otherwise submit or reference to support an application, are not sufficient to support approval of a product candidate; the FDA or comparable regulatory authorities in other countries may ultimately conclude that our manufacturing processes or facilities or those of our third-party manufacturers do not sufficiently demonstrate compliance with cGMP to support approval of a product candidate; that the drug CMC data or device biocompatibility data for our product candidates otherwise do not support approval or that additional CMC data or information for our product candidates must be submitted for review; 63Table of Contents we may be unable to demonstrate to the satisfaction of the FDA or comparable regulatory authorities in other countries that our product candidate is safe and effective for its proposed indication, or that its clinical and other benefits outweigh its safety risks; the approval policies of the FDA or comparable regulatory authorities in other countries may change in a manner that renders our data insufficient for approval. Even if we obtain marketing approval, the FDA or comparable regulatory authorities in other countries may approve our product candidates for fewer or more limited indications than those for which we requested approval or may include safety warnings or other restrictions that may negatively impact the commercial viability of our product candidates. Likewise, regulatory authorities may grant approval contingent on the performance of costly post-marketing clinical trials or other studies or the conduct of an expensive risk evaluation and mitigation strategies, or REMS, which could significantly reduce the potential for commercial success or viability of our product candidates. We also may not be able to find acceptable collaborators to manufacture our drug products, if and when approved, in commercial quantities and at acceptable prices, or at all. We may encounter difficulties in enrolling patients in our clinical trials. We may not be able to commence or complete clinical trials for our product candidates if we are unable to locate and enroll a sufficient number of eligible patients to participate in these trials. Patient enrollment may be affected by a variety of factors, including, among others: the severity of the disease under investigation; the design of the clinical trial protocol and amendments to a protocol; the size and nature of the patient population; eligibility criteria for the clinical trial in question; the perceived risks and benefits of the product candidate under clinical testing, including a high and unacceptable severity, or prevalence, of undesirable side effects or adverse events caused by our product candidates or similar products or product candidates; the existing body of safety and efficacy data in respect of the product candidate under clinical testing; the proximity of patients to clinical trial sites; the number and nature of competing therapies and clinical trials; and other environmental factors such as pandemics or other natural or unforeseen disasters. Any negative results we may report in clinical trials of our product candidates may also make it difficult or impossible to recruit and retain patients in other clinical trials of that same product candidate. We expect that if we initiate, as we are currently contemplating, a clinical trial of YUTREPIA in pediatric patients, we may encounter difficulties enrolling patients in such a trial because of the limited number of pediatric patients with this disease. Furthermore, we are aware of a number of therapies for PAH that are being developed or that are already available on the market, and we expect to face competition from these investigational drugs or approved drugs for potential subjects in our clinical trials, including planned clinical trials for YUTREPIA and L606, which may delay enrollment in our planned clinical trials. Delays or failures in planned patient enrollment or retention may result in increased costs, program delays, or both. We may, as a result of such delays or failures, be unable to carry out our clinical trials as planned or within the timeframe that we expect or at all, and our business and prospects may be materially and adversely affected as a result. 64Table of Contents Product candidates that the FDA deems to be combination products, such as YUTREPIA and L606, or that otherwise rely on innovative drug delivery systems, may face additional challenges, risks and delays in the product development and regulatory approval process. The FDA has indicated that it considers YUTREPIA, which is delivered by a DPI, and L606, which is delivered by a next generation nebulizer, to be drug-device combination products. Accordingly, the medical devices used to administer the products were, or in the case of L606 will be, evaluated as part of our NDA filing. When evaluating products that utilize a specific drug delivery system or device, the FDA will evaluate the characteristics of that delivery system and its functionality, as well as the potential for undesirable interactions between the drug and the delivery system, including the potential to negatively impact the safety or effectiveness of the drug. The FDA review process can be more complicated for combination products, and may result in delays, particularly if novel delivery systems are involved. We rely on third parties for the

design and manufacture of the delivery systems for our products, including the DPI for YUTREPIA and the nebulizer for L606, and in some cases for the right to refer to their data on file with the FDA or other regulators. Quality or design concerns with the delivery system, or commercial disputes with these third parties, could delay or prevent regulatory approval and commercialization of our product candidates. We are pursuing the FDA 505(b)(2) pathway for our current product candidates. If we are unable to rely on the 505(b)(2) regulatory pathway to apply for marketing approval of our product candidates in the United States, seeking approval of these product candidates through the 505(b)(1) NDA pathway would require full reports of investigations of safety and effectiveness, and the process of obtaining marketing approval for our product candidates would likely be significantly longer and more costly. We are currently focused on developing drug products that can be approved under abbreviated regulatory pathways in the United States, such as the 505(b)(2) regulatory pathway, which permits the filing of an NDA where at least some of the information required for approval comes from studies that were not conducted by or for the applicant and for which the applicant has not obtained a right of reference. Section 505(b)(2), if applicable to us for a particular product candidate, would allow an NDA we submit to the FDA to rely in part on data in the public domain or the FDA's prior conclusions regarding the safety and effectiveness of approved compounds, which could expedite the development program for a product candidate by potentially decreasing the amount of clinical data that we would need to generate in order to obtain FDA approval. We have pursued this pathway for our current product candidate, YUTREPIA, and are pursuing this pathway for L606. Even if the FDA allows us to rely on the 505(b)(2) regulatory pathway for a given product candidate, we cannot assure you that marketing approval will be obtained in a timely manner, or at all. The FDA may require us to perform additional clinical trials to support any change from the reference listed drug, which could be time-consuming and substantially delay our receipt of marketing approval. Also, as has been the experience of others in our industry, our competitors may file citizen petitions or other correspondence with the FDA or lawsuits against the FDA to contest approval of our NDA, which may delay or even prevent the FDA from approving any NDA that we submit under the 505(b)(2) regulatory pathway. For instance, United Therapeutics has a lawsuit against the FDA and recently filed a citizen petition in an attempt to prevent or delay the approval of YUTREPIA. If an FDA decision or action relative to our product candidate, or the FDA's interpretation of Section 505(b)(2) more generally, is successfully challenged, it could result in delays or even prevent the FDA from approving a 505(b)(2) application for our product candidates or for certain indications for our product candidates. Even if we are able to utilize the 505(b)(2) regulatory pathway, the approval of a drug developed under the 505(b)(2) regulatory pathway may be delayed by one or more regulatory exclusivities. For example, Tyvaso DPI was recently granted New Clinical Investigation exclusivity, which has delayed final approval of YUTREPIA until after the exclusivity expires in May 2025. Also, a drug approved via this pathway may be subject to the same post-approval limitations, conditions and requirements as any other drug. In addition, we may face Hatch-Waxman litigation in relation to our NDAs submitted under the 505(b)(2) regulatory pathway, which may further delay or prevent the approval of our product candidates. The pharmaceutical industry is highly competitive, and 505(b)(2) NDAs are subject to special requirements designed to protect the patent rights of sponsors of previously approved drugs that are referenced in a 505(b)(2) NDA. If the previously approved drugs referenced in an applicant's 505(b)(2) NDA are protected by patent(s) listed in the Orange Book, the 505(b)(2) applicant is required to make a claim after filing its NDA or certain types of amendments to its NDA that each such patent is invalid, unenforceable or will not be infringed. The patent holder may thereafter bring suit for patent infringement, 65Table of Contentswhich will trigger a mandatory 30-month delay (or the shorter of dismissal of the lawsuit or expiration of the patent(s)) in approval of the 505(b)(2) NDA application. In addition, in the event the court in any such lawsuit finds that any claims of any of the asserted patents are both valid and infringed, the court would likely issue an injunction prohibiting approval of the product at issue until the expiration of the patent(s) found to have been infringed. For example, the YUTREPIA NDA was filed under the 505(b)(2) regulatory pathway with Tyvaso as the reference listed drug. Under the Hatch-Waxman Act, as a result of the litigation commenced by United Therapeutics in June 2020, the FDA was automatically precluded from approving the YUTREPIA NDA for up to 30 months. Also, in connection with an amendment to our NDA filed in July 2023 to add PH-ILD as an indication for YUTREPIA, we provided a new notice of the paragraph IV certification to United Therapeutics as the owner of the patents that are the subject of the certification to which the NDA for YUTREPIA refers. As a result, in September 2023, United Therapeutics filed the New Hatch-Waxman Litigation, again asserting infringement by the Company of the '793 Patent, which lawsuit was amended on November 30, 2023, to add claims asserting infringement of the '327 Patent. In February 2024, United Therapeutics filed a motion seeking a preliminary injunction to prevent us from manufacturing, marketing, storing, importing, distributing, offering for sale, and/or selling YUTREPIA for the treatment of PH-ILD. Although the motion for preliminary injunction was denied, United Therapeutics may still seek injunctive relief and other remedies. In addition, United Therapeutics may seek to assert newly issued patents against us, including U.S. Patent Number 11,723,887, and may seek to enjoin the FDA from granting final approval to YUTREPIA or enjoin us from launching YUTREPIA. It is also not uncommon for a manufacturer of an approved product, such as United Therapeutics, to file a citizen petition or other correspondence with the FDA seeking to delay approval of, or impose additional approval requirements for, pending competing products or to take other actions, such as engaging in litigation with the FDA to enjoin approval of a competing product. If successful, such petitions, correspondence or litigation can significantly delay, or even prevent, the approval of the new product. For example, United Therapeutics is currently pursuing litigation under the Administrative Procedures Act, seeking to require the FDA to reject our amendment to the YUTREPIA NDA to add PH-ILD to the label. Even if the FDA ultimately prevails in such litigation, the FDA may substantially delay approval while it considers and responds to the petition or correspondence and is engaged in litigation or the FDA may be temporarily enjoined by a court from granting approval until the court has ruled on United Therapeutics' request. If the FDA determines that any of our product candidates do not qualify for the 505(b)(2) regulatory pathway, we would need to reconsider our plans and might not be able to commercialize our product candidates in a cost-efficient manner, or at all. If we were to pursue approval under the 505(b)(1) NDA pathway, we would be subject to more extensive requirements and risks such as conducting additional clinical trials, providing additional data and information or meeting additional standards for marketing approval. As a result, the time and financial resources required to obtain marketing approval for our product candidates would likely increase substantially and further complications and risks associated with our product candidates may arise. Also, new competing products may reach the market faster than ours, which may materially and adversely affect our competitive position, business and prospects. We may be unable to continually develop a pipeline of product candidates, which could affect our business and prospects. A key element of our long-term strategy is to continually develop a pipeline of product candidates by developing products for the treatment of pulmonary hypertension and proprietary innovations to FDA-approved drug products using our PRINT technology. If we are unable to identify suitable product candidates for the treatment of pulmonary hypertension or off-patent drug products for which we can develop proprietary innovations using our PRINT technology or are otherwise unable to expand our product candidate pipeline, whether through licensed or co-development opportunities, and obtain marketing approval for such product candidates within the timeframes that we anticipate, or at all, our business and prospects may be materially and adversely affected. 66Table of ContentsWe have conducted, and may in the future conduct, clinical trials for our product candidates outside the United States and the FDA may not accept data from such trials. Although the FDA may accept data from clinical trials conducted outside the United States in support of safety and efficacy claims for our product candidates, if not conducted under an IND, this is subject to certain conditions set out in 21 C.F.R. § 312.120. For example, in order for the FDA to accept data from such a foreign clinical trial, the study must have been conducted in accordance with Good Clinical Practice (GCP) including review and approval by an independent ethics committee and obtaining the informed consent from subjects of the clinical trials. The FDA must also be able to validate the data from the study through an onsite inspection if the agency deems it necessary. In addition, foreign clinical data submitted to support FDA applications should be applicable to the U.S. population and U.S. medical practice. Other factors that may affect the acceptance of foreign clinical data include differences in clinical conditions, study populations or regulatory requirements between the United States and the foreign country. Risks Related to Our Dependence on Third PartiesWe depend on third parties for clinical and commercial supplies, including single suppliers for the active ingredient, the device, encapsulation and packaging of YUTREPIA and single suppliers for the active ingredient, bulk product manufacturing and packaging of L606. We depend on third-party suppliers for clinical and commercial supplies for the supply of materials and components necessary for clinical and commercial production of YUTREPIA and L606, including the active pharmaceutical ingredients which are used in our product candidates. These supplies may not always be available to us at the standards we require or on terms acceptable to us, or at all, and we may not be able to locate alternative suppliers in a timely manner, or at all. If we are unable to obtain necessary clinical or commercial supplies, our manufacturing operations and clinical trials and the clinical trials of our collaborators may be delayed or disrupted and our business and prospects may be materially and adversely affected as a result. For example, we currently rely on a sole supplier for treprostinil, the active pharmaceutical ingredient of YUTREPIA, which sources treprostinil from a manufacturer in South Korea, with whom we have a long-term supply agreement. If our supplier is unable to supply treprostinil to us in the quantities we require, or at all, or otherwise defaults on its supply obligations to us, or if it ceases its relationship with us, we may not be able to obtain alternative supplies of treprostinil from other suppliers on acceptable terms, in a timely manner, or at all. We also rely on a sole supplier located in Tampa, Florida for encapsulation and packaging services, with whom we have a long-term contract. Furthermore, YUTREPIA is administered using the RS00 Model 8 DPI, which is manufactured by Plastiapi, which is located in Italy. In the event of any prolonged disruption to our supply of treprostinil, the encapsulation and packaging services, or the manufacture and supply of RS00 Model 8 DPI, our ability to develop and commercialize, and the timeline for commercialization of, YUTREPIA may be adversely affected. We also rely upon Chengdu for the manufacture and supply of RG Cartridges for the subcutaneous administration of Treprostinil Injection and upon ICU Medical for ongoing servicing and support of the CADD-MS 3, CADD Legacy and CADD-Solis infusion pumps. In the event of any disruption to our supply of RG Cartridges or any disruption in the availability of parts or servicing for the CADD-MS 3, CADD Legacy and CADD-Solis infusion pumps, sales of Treprostinil Injection may be adversely affected. In addition, ICU Medical has indicated that they will no longer support the CADD-MS 3. Although we believe that the number of available CADD-MS 3 infusion pumps will be sufficient to continue serving patients through the end of 2025, we are relying upon Mainbridge for the development of new pumps for the subcutaneous administration of Treprostinil Injection to replace the CADD-MS-3. In addition, we have still not submitted a 510(k) clearance application and are currently uncertain when, if ever, such a 510(k) clearance application will be submitted. If we are unable to identify options to maintain the availability of the existing CADD-MS-3 pumps until the new pumps are cleared by the FDA, sales of Treprostinil Injection may be adversely affected. For L606, we rely upon single sources of supply for the active pharmaceutical ingredient, manufacture of bulk drug product and packaging. Some of these suppliers are located in Taiwan. Although we are working to establish a 67Table of Contentssecondary supply chain outside of Taiwan, if hostilities were to break out between Taiwan and China, we may be unable to secure a supply of L606. Also, we are currently evaluating devices to use for the administration of L606. If we are unable to identify a device to use for our L606 program, establish an agreement with the manufacturer of that device for the supply of such devices or obtain adequate quantities of that device in a timely manner or at all, we may be unable to successfully develop L606 or to do so in a timely manner. If any of our sole source suppliers are adversely affected by geopolitical events, natural disasters or other events that disrupt or adversely affect their operations or their ability to supply us, our business may be adversely affected. If we are unable to establish or maintain licensing and collaboration arrangements with other pharmaceutical companies on acceptable terms, or at all, we may not be able to develop and commercialize additional product candidates using our PRINT technology. We have collaborated, and may consider collaborating, with, among others, pharmaceutical companies to expand the applications for our PRINT technology through licensing as well as joint product development arrangements. In addition, if we are able to obtain marketing approval for our product candidates from regulatory authorities, we may enter into strategic relationships with collaborators for the commercialization of such products. Collaboration and licensing arrangements are complex and time-consuming to negotiate, document, implement and maintain. We may not be successful in our efforts to establish collaboration or other alternative arrangements should we so choose to enter into such arrangements. In addition, the terms of any collaboration or other arrangements that we may enter into may not be favorable to us or may restrict our ability to enter into further collaboration or other arrangements with third parties. For example, collaboration agreements may contain exclusivity arrangements which limit our ability to work with other pharmaceutical companies to expand the applications for our PRINT technology, as is the case in our collaboration agreement with GSK which restricts our ability to use PRINT for inhaled applications with respect to certain identified compounds. If we are unable to establish licensing and collaboration arrangements on the terms of such agreements we enter into are unfavorable to us or restrict our ability to work with other pharmaceutical companies, we may not be able to expand the applications for our PRINT technology or commercialize our products, if and when approved, and our business and prospects may be materially and adversely affected. Our collaboration and licensing arrangements may not be successful. Our collaboration and licensing arrangements, as well as any future collaboration and licensing arrangements that we may enter into, may not be successful. The success of our collaboration and licensing arrangements will depend heavily on the efforts and activities of our collaborators, which are not within our control. We may, in the course of our collaboration and licensing arrangements, be subject to numerous risks, including, but not limited to, the following: our collaborators may have significant discretion in determining the efforts and resources that they will contribute; our collaborators may delay clinical trials, provide insufficient funding for a clinical trial program, stop a clinical trial, abandon a product candidate, repeat or conduct new clinical trials or require a new formulation of a product candidate for clinical testing. For example, in July 2018, GSK notified us of its decision to discontinue development of the inhaled antiviral for viral exacerbations in COPD after completion of its related Phase 1 clinical trial and we do not believe that GSK is currently advancing any program under our collaboration; our collaborators may independently, or in conjunction with others, develop products that compete directly or indirectly with our product candidates; we may grant exclusive rights to our collaborators that would restrict us from collaborating with others. For example, we are currently subject to certain restrictions with regard to our ability to enter into collaboration arrangements to use PRINT for the development of inhaled therapeutics using certain identified compounds pursuant to our collaboration with GSK; our collaborators may not properly maintain or defend our intellectual property rights or may use our intellectual property or proprietary information in a way that gives rise to actual or threatened litigation that could jeopardize or invalidate our intellectual property or proprietary information or expose us to potential liability; disputes may arise between us and our collaborators, which may cause a delay in or the termination of our research, development or commercialization activities; our collaboration and licensing arrangements may be terminated, and if terminated, may result in our need for additional capital to pursue further drug product development or commercialization. For example, our development and licensing agreement with G&W Laboratories, Inc., was mutually terminated in April 2018; our collaborators may own or co-own certain intellectual property arising from our collaboration and licensing arrangements with them, which may restrict our ability to develop or commercialize such intellectual property; and our collaborators may alter the strategic direction of their business or may undergo a change of control or management, which may affect the success of our collaboration arrangements with them. Risks Related to our Intellectual PropertyWe may be subject to claims from third parties that our products infringe their intellectual property rights. The pharmaceutical industry has experienced rapid technological change and obsolescence in the past, and our competitors have strong incentives to stop or delay any introduction of new drug products or related technologies by, among others, establishing intellectual property rights over their drug products or technologies and aggressively enforcing these rights against potential new entrants into the market. We expect that we and other industry participants will be increasingly subject to infringement claims as the number of competitors and drug products grows. Our commercial success depends in large part upon our ability to develop, manufacture, market and sell our drug products or product candidates without infringing on the patents or other proprietary rights of third parties. It is not always clear to industry participants, including us, what the scope of a patent covers. Due to the large number of patents in issue and patent applications filed in our industry, there is a risk that third parties will claim that our products or technologies infringe their intellectual property rights. Claims for infringement of intellectual property which are brought against us, whether with or without merit, and which are generally uninsurable, could result in time-consuming and costly litigation, diverting our management's attention from our core business and reducing the resources available for our drug product development, manufacturing and marketing activities, and consequently have a material and adverse effect on our business and prospects, regardless of the outcome. Moreover, such proceedings could put our patents at risk of being invalidated or interpreted narrowly and our patent applications at risk of not being issued. We also may not prevail in any lawsuits that we initiate and the damages or other remedies awarded, if any, may not be commercially meaningful. Uncertainties resulting from the initiation and continuation of litigation or other proceedings could also have a material and adverse effect on our ability to compete in the market. Third parties making claims against us could obtain injunctive or other equitable relief against us, which could prevent us from further developing or commercializing our product candidates. In particular, under the Hatch-Waxman Act, the owner of patents listed on the Orange Book and referenced by an NDA applicant may bring patent infringement suit against the NDA applicant after receipt of the NDA applicant's notice of paragraph IV certification. For example, in connection with an amendment to our NDA filed in July 2023 to add PH-ILD as an indication for YUTREPIA, a new

notice of the paragraph IV certification was provided to United Therapeutics as the owner of the patents that are the subject of the certification to which the NDA for YUTREPIA refers. As a result, United Therapeutics filed the New Hatch-Waxman Litigation, in which it sought a preliminary injunction. While the motion for a preliminary injunction was denied, United Therapeutics may still seek injunctive relief in the New Hatch-Waxman Litigation. Although we do not believe United Therapeutics is entitled to a preliminary injunction in connection with the New Hatch-Waxman Litigation, it is possible that the Court could enjoin us from commercializing YUTREPIA for the treatment of PH-ILD. ^{69Table of Contents}In addition, United Therapeutics may seek to assert newly issued patents against us, including U.S. Patent Number 11,723,887, and may seek to enjoin the FDA from granting final approval to YUTREPIA or enjoin us from launching YUTREPIA, including through temporary restraining orders or injunctions that they may seek in the New FDA Litigation.⁶⁹In the event of a successful infringement claim against us, including an infringement claim filed in response to a paragraph IV certification, we may be required to pay damages, cease the development or commercialization of our drug products or product candidates, limit the label of our products to fewer indications than intended, re-engineer or redevelop our drug products or product candidates or enter into royalty or licensing agreements, any of which could have a material and adverse impact on our business, financial condition and results of operations. Any effort to re-engineer or redevelop our products would require additional monies and time to be expended and may not ultimately be successful.⁶⁹Infringement claims may be brought against us in the future, and we cannot assure you that we will prevail in any ensuing litigation given the complex technical issues and inherent uncertainties involved in intellectual property litigation. Our competitors may have substantially greater resources than we do and may be able to sustain the costs of such litigation more effectively than we can.⁶⁹Our commercial success depends largely on our ability to protect our intellectual property. Our commercial success depends, in large part, on our ability to obtain and maintain patent protection and trade secret protection in the United States and elsewhere in respect of our product candidates and PRINT technology. If we fail to adequately protect our intellectual property rights, our competitors may be able to erode, negate or preempt any competitive advantage we may have. To protect our competitive position, we have filed and will continue to file for patents in the United States and elsewhere in respect of our product candidates and PRINT technology. The process of identifying patentable subject matter and filing a patent application is expensive and time-consuming. We cannot assure you that we will be able to file the necessary or desirable patent applications at a reasonable cost, in a timely manner, or at all. Further, since certain patent applications are confidential until patents are issued, third parties may have filed patent applications for subject matter covered by our pending patent applications without us being aware of such applications, and our patent applications may not have priority over patent applications of others. In addition, we cannot assure you that our pending patent applications will result in patents being obtained. Once published, all patent applications and publications throughout the world, including our own, become prior art to our new patent applications and may prevent patents from being obtained or interfere with the scope of patent protection that might be obtained. The standards that patent offices in different jurisdictions use to grant patents are not always applied predictably or uniformly and may change from time to time.⁶⁹Even if we have been or are able to obtain patent protection for our product candidates or PRINT technology, if the scope of such patent protection is not sufficiently broad, we may not be able to rely on such patent protection to prevent third parties from developing or commercializing product candidates or technology that may copy our product candidates or technology. The enforceability of patents in the pharmaceutical industry involves complex legal and scientific questions and can be uncertain. Accordingly, we cannot assure you that third parties will not successfully challenge the validity, enforceability or scope of our patents. A successful challenge to our patents may lead to generic versions of our drug products being launched before the expiry of our patents or otherwise limit our ability to stop others from using or commercializing similar or identical products and technology. A successful challenge to our patents may also reduce the duration of the patent protection of our drug products or technology. In addition, we cannot assure you that we will be able to detect unauthorized use or take appropriate, adequate and timely actions to enforce our intellectual property rights. If we are unable to adequately protect our intellectual property, our business, competitive position and prospects may be materially and adversely affected.⁶⁹Even if our patents or patent applications are unchallenged, they may not adequately protect our intellectual property or prevent third parties from designing around our patents or other intellectual property rights. If the patent applications we file or may file do not lead to patents being granted or if the scope of any of our patent applications is challenged, we may face difficulties in developing our product candidates, companies may be dissuaded from collaborating with us, and our ability to commercialize our product candidates may be materially and adversely affected. We are unable to predict which of our patent applications will lead to patents or assure you that any of our patents will not be found invalid or ^{70Table of Contents}unenforceable or challenged by third parties. The patents of others may prevent the commercialization of product candidates incorporating our technology. In addition, given the amount of time required for the development, clinical testing and regulatory review of new product candidates, any patents protecting our product candidates may expire before or shortly after such product candidates might become approved for commercialization.⁶⁹Moreover, the issuance of a patent is not conclusive as to the inventorship of the patented subject matter, or its scope, validity or enforceability. We cannot assure you that all of the potentially relevant prior art, that is, any evidence that an invention is already known, relating to our patents and patent applications, has been found. If such prior art exists, it may be used to invalidate a patent or may prevent a patent from being issued.⁶⁹Questions may also arise as to the ownership of our patents. For instance, in May 2024, United Therapeutics filed a complaint in the Superior Court in Durham County, North Carolina, in which it is seeking declaratory judgement such that all right, title and interest in and to any patentable or unpatentable inventions, discoveries, and ideas made or conceived by Dr. Roscigno while employed by the Company should be assigned and transferred to United Therapeutics because they involved the use of United Therapeutics' ⁷¹confidential information. If successful, United Therapeutics could obtain an ownership interest in our patents, which may either limit our ability to prevent United Therapeutics from using out-patented inventions or even allow United Therapeutics to prevent us from using our own patented inventions.⁶⁹In addition, we, our collaborators or our licensees may fail to identify patentable aspects of inventions made in the course of development and commercialization activities before it is too late to obtain patent protection on them. As a result, we may miss potential opportunities to seek patent protection or strengthen our patent position.⁶⁹If we are unable to protect our trade secrets, the value of our PRINT technology and product candidates may be negatively impacted, which would have a material and adverse effect on our competitive position and prospects. In addition to patent protection, we rely on trade secret protection to protect certain aspects of our intellectual property. We also license trade secrets from Pharmosa with respect to L606. While we require parties who have access to any portion of our trade secrets, such as our employees, consultants, advisers, CROs, CMOs, collaborators and other third parties, to enter into non-disclosure and confidentiality agreements with us, we cannot assure you that these parties will not disclose our proprietary information, including our trade secrets, in breach of their contractual obligations. Enforcing a claim that a party has illegally disclosed or misappropriated a trade secret is difficult, costly and time-consuming, and we may not be successful in doing so. If the steps we have taken to protect our trade secrets are deemed by the adjudicating court to be inadequate, we may not be able to obtain adequate recourse against a party for misappropriating our trade secrets.⁶⁹Trade secrets can be difficult to protect as they may, over time, be independently discovered by our competitors or otherwise become known despite our trade secret protection. If any of our trade secrets were to be lawfully obtained or independently developed by our competitors, we would have no right to prevent such competitors, or those to whom they communicate such technology or information, from using that technology or information to compete with us. Such competitors could attempt to replicate some or all of the competitive advantages we derive from our development efforts, willfully infringe our intellectual property rights, design around our protected technology or develop their own competitive technologies that fall outside of our intellectual property rights.⁶⁹If our trade secrets were to be disclosed to or independently developed by our competitors, our competitors may be able to exploit our PRINT technology to develop competing product candidates, and the value of our PRINT technology and our product candidates may be negatively impacted. This would have a material and adverse effect on our competitive position and prospects.⁶⁹We rely on licenses to intellectual property that are owned by third parties. We have entered and may, in the future, enter into license agreements with third parties to license the rights to use their technologies in our research, development and commercialization activities. License agreements generally impose various diligence, milestone payment, royalty, insurance and other obligations on us, and if we fail to comply with these obligations, our licensors may have the right to terminate these license agreements. Termination of these license ^{71Table of Contents}agreements or the reduction or elimination of our licensed rights or the exclusivity of our licensed rights may have an adverse impact on, among others, our ability to develop and commercialize our product candidates. We cannot assure you that we will be able to negotiate new or reinstated licenses on commercially acceptable terms, or at all.⁶⁹In addition, we license certain patent rights for our PRINT technology from UNC under the UNC License. Under the UNC License, UNC has the right to terminate our license if we materially breach the agreement and fail to cure such breach within the stipulated time. In the event that UNC terminates our license and we have a product that relies on that license, including YUTREPIA, it may bring a claim against us, and if they are successful, we may be required to compensate UNC for the unauthorized use of their patent rights through the payment of royalties.⁶⁹Similarly, under our license agreement with Pharmosa, Pharmosa has the right to terminate our license if we materially breach the agreement and fail to cure such breach within the stipulated time. In the event that Pharmosa terminates our license and we have a product that relies on that license, including L606, it may bring a claim against us, and if they are successful, we may be required to compensate Pharmosa for the unauthorized use of their patent rights through the payment of royalties.⁶⁹Also, the agreements under which we license patent rights may not give us control over patent prosecution or maintenance, so that we may not be able to control which claims or arguments are presented and may not be able to secure, maintain or successfully enforce necessary or desirable patent protection from those patent rights. We do not have primary control over patent prosecution and maintenance for certain of the patents we license, and therefore cannot assure you that these patents and applications will be prosecuted or maintained in a manner consistent with the best interests of our business. We also cannot assure you that patent prosecution and maintenance activities by our licensors, if any, will be conducted in compliance with applicable laws and regulations or will result in valid and enforceable patents.⁶⁹Pursuant to the terms of some of our license agreements with third parties, some of our third-party licensors have the right, but not the obligation, in certain circumstances, to control the enforcement of our licensed patents or defense of any claims asserting the invalidity of these patents. Even if we are permitted to pursue such enforcement or defense, we will require the cooperation of our licensors, and we cannot assure you that we will receive such cooperation on commercially acceptable terms, or at all. We also cannot assure you that our licensors will allocate sufficient resources or prioritize their or our enforcement of these patents or defense of these claims to protect our interests in the licensed patents. If we cannot obtain patent protection, or enforce existing or future patents against third parties, our competitive position, business and prospects may be materially and adversely affected.⁶⁹Further, licenses to intellectual property may not always be available to us on commercially acceptable terms, or at all. In the event that the licenses we rely on are not available to us on commercially acceptable terms, or at all, our ability to commercialize our PRINT technology or product candidates, and our business and prospects, may be materially and adversely affected.⁶⁹We may not be able to enforce our intellectual property rights throughout the world. Filing, prosecuting, enforcing and defending patents on our PRINT technology and our product candidates throughout the world may be prohibitively expensive and may not be financially or commercially feasible. In countries where we have not obtained patent protection, our competitors may be able to use our proprietary technologies to develop competing product candidates.⁶⁹Also, the legal systems of non-U.S. jurisdictions may not protect intellectual property rights to the same extent or in the same manner as the laws of the United States, and we may face significant difficulty in enforcing our intellectual property rights in these jurisdictions. The legal systems of certain developing countries may not favor the enforcement of patents and other intellectual property rights. We may therefore face difficulty in stopping the infringement or misappropriation of our patents or other intellectual property rights in those countries.⁶⁹^{72Table of Contents}We need to protect our trademark, trade name and service mark rights to prevent competitors from taking advantage of our name recognition. We believe that the protection of our trademark, trade name and service mark rights, such as Liquidia, the Liquidia logo, PRINT, and YUTREPIA, is an important factor in product recognition, protecting our brand, maintaining goodwill and maintaining or increasing market share. We may expend substantial cost and effort in an attempt to register new trademarks, trade names and service marks and maintain and enforce our trademark, trade name and service mark rights. If we do not adequately protect our rights in our trademarks, trade names and service marks from infringement, any name recognition that we have developed in those trademarks could be lost or impaired.⁶⁹Third parties may claim that the sale or promotion of our products, when and if approved, may infringe on the trademark, trade name and service mark rights of others. Trademark, trade name and service mark infringement problems occur frequently in connection with the sale and marketing of pharmaceutical products. If we become involved in any dispute regarding our trademark, trade name and service mark rights, regardless of whether we prevail, we could be required to engage in costly, distracting and time-consuming litigation that could harm our business. If the trademarks, trade names and service marks we use are found to infringe upon the trademarks, trade names or service marks of another company, we could be liable for damages and be forced to stop using those trademarks, trade names or service marks, and as a result, we could lose all the name recognition that has been developed in those trademarks, trade names or service marks.⁶⁹Risks Related to the Manufacturing of our Product Candidates⁶⁹Our product candidates are based on our proprietary, novel technology, which has not been used to manufacture any products that have been previously approved by the FDA, making it difficult to predict the time and cost of development and of subsequently obtaining final regulatory approval. Our future success depends on the successful development of our novel PRINT technology and products based on it, including YUTREPIA, and the development of L606 using Pharmosa's proprietary liposomal technology. To our knowledge, no regulatory authority has granted final approval to market or commercialize drugs made using our PRINT technology or Pharmosa's liposomal technology. We may never receive final approval to market and commercialize any product candidate that uses our PRINT technology or Pharmosa's liposomal technology. Even if we receive final approval to market YUTREPIA and/or L606, we will need to scale up our manufacturing capabilities to effectively commercialize the products. We have never completed a scale up of our PRINT manufacturing process or the manufacturing process for L606, and, if we are unable to do so in an effective and timely manner, our ability to commercialize these products, even if they receive final FDA approval, will be adversely affected. We may experience unexpected challenges as we ramp up our manufacturing capacity to meet demand or during commercial manufacturing, which may result in our inability to supply sufficient quantities of product to meet demand. The manufacturing process for our products is complex, due in part to strict regulatory requirements. A failure of our quality control systems in our facilities or those of our CMOs could cause problems to arise in connection with facility operations for a variety of reasons, including equipment malfunction, viral contamination, failure to follow specific manufacturing instructions, protocols and standard operating procedures, problems with raw materials or environmental factors. Such problems could affect production of a single batch or a series of batches, requiring the destruction of products, or could halt manufacturing operations altogether. For instance, as we scale up the manufacture of YUTREPIA, we are adjusting the speed and temperature at which our blister packs are sealed to reduce the risk of the product being exposed to moisture. Our failure to meet required quality standards may result in our failure to timely deliver products to our customers in sufficient quantities to meet demand, which in turn could damage our reputation for quality and service. Any such incident could, among other things, lead to increased costs, lost revenue, damage to our reputation and relationships with patients, health care providers and third-party payors, time and expense spent investigating the cause and, depending on the cause, similar losses with respect to other batches. With respect to our commercial manufacturing, if problems are not discovered before the product is released to the market, we may be ^{73Table of Contents}subject to regulatory actions, including product recalls, product seizures, injunctions to halt manufacture and distribution, restrictions on our operations, civil sanctions, including monetary sanctions, and criminal actions. In addition, such issues could subject us to litigation, the cost of which could be significant. Our operations are concentrated in Morrisville, North Carolina and interruptions affecting us or our suppliers due to natural disasters or other unforeseen events could materially and adversely affect our operations. Most of our current operations are concentrated in Morrisville, North Carolina. In addition, our inventory is warehoused in a limited number of locations. A fire, flood, hurricane, earthquake or other disaster or unforeseen event resulting in significant damage to our facilities or to inventory held by us could significantly disrupt or curtail or require us to cease our operations. It would be difficult, costly and time-consuming to transfer resources from one facility to another, to repair or replace our facility or to replace inventory in the event that it is significantly damaged. In addition, our insurance may not be sufficient to cover all of our losses and may not continue to be available to us on acceptable terms, or at all. In addition, if one of our suppliers experiences a similar disaster or unforeseen event, we could face significant loss of our inventory and significant delays in obtaining our supplies or be required to source supplies from an alternative supplier and may incur substantial costs as a result. Any significant uninsured loss, prolonged or repeated disruption to operations or inability to operate, experienced by us or by our suppliers, could materially and adversely affect our business, financial condition and results of operations.⁶⁹In addition, for L606, we rely upon single sources of supply for the active pharmaceutical ingredient and manufacture of bulk drug that are located in Taiwan. Although we are working to establish a secondary supply chain outside of Taiwan, if hostilities were to break out between Taiwan and China, we may be unable to secure a supply of L606, which could limit our ability to continue development of L606 and materially

and adversely affect our business, financial condition and results of operations. Risks Related to our Employees We depend on skilled labor, and our business and prospects may be adversely affected if we lose the services of our skilled personnel, including those in senior management, or are unable to attract new skilled personnel. Our ability to continue our operations and manage our potential future growth depends on our ability to hire and retain suitably skilled and qualified employees, including those in senior management, in the long-term. Due to the specialized nature of our work, there is a limited supply of suitable candidates. We compete with other biotechnology and pharmaceutical companies, educational and research institutions and government entities, among others, for research, technical, clinical and sales and marketing personnel. In addition, in order to manage our potential future growth effectively, we will need to improve our financial controls and systems and, as necessary, recruit sales, marketing, managerial and finance personnel. The loss of the services of members of our sales team could seriously harm our ability to successfully implement our business strategy. If we are unable to attract and retain skilled personnel, including in particular Roger Jeffs, our Chief Executive Officer, we may not be able to successfully implement the tasks necessary to further develop and commercialize our product candidates and, accordingly, our business and prospects may be materially and adversely affected. Risks Related to our Common Stock Future sales of our Common Stock or securities convertible into our Common Stock in the public market could cause our stock price to fall. Our stock price could decline as a result of sales of a large number of shares of our Common Stock or the perception that these sales could occur. These sales, or the possibility that these sales may occur, also might make it more difficult for us to sell equity securities in the future at a time and at a price that we deem appropriate. As of October 30, 2024, 84,636,621 shares of our Common Stock were outstanding, of which 76,405,243 shares of Common Stock, or 90.3% of our outstanding shares as of October 30, 2024, are freely tradable without restriction or further registration under the Securities Act, provided however, some of these shares are held by persons deemed to be 74Table of Contentsâ€œaffiliatesâ€ under the Securities Act, including our officers and directors, as well as our principal stockholders, and may not be sold except: (i) in compliance with Rule 144 under the Securities Act or (ii) pursuant to any other applicable exemption under the Securities Act. The remaining 8,231,378 shares held by our stockholders as of October 30, 2024 have not been registered under the Securities Act and may be only be sold (i) pursuant to an effective registration statement under the Securities Act covering the sale of those shares, (ii) in compliance with Rule 144 under the Securities Act or (iii) pursuant to any other applicable exemption under the Securities Act. Shares issued upon purchase under the employee stock purchase plan or upon the exercise of stock options or vesting of restricted stock units outstanding under our equity incentive plans or pursuant to future awards granted under those plans will become available for sale in the public market to the extent permitted by the provisions of applicable vesting schedules, any applicable market standoff and lock-up agreements, and Rule 144 and Rule 701 under the Securities Act. We have registered the offer and sale of all shares of Common Stock that we may issue under our equity compensation plans, including the employee stock purchase plan. We expect that the market price of our Common Stock may be volatile, and you may lose all or part of your investment. The trading prices of the securities of pharmaceutical and biotechnology companies have been highly volatile. As such, the trading price of our Common Stock may be highly volatile and could be subject to wide fluctuations in response to various factors, some of which are beyond our control. The market price for our Common Stock may be influenced by many factors, including:â€—results of any clinical trials of any product candidate we may develop, including L606, or those of our competitors;â€—the success of Sandozâ€™s Troprelin Injection to which we have commercial rights pursuant to the Promotion Agreement;â€—the market acceptance of the RG Cartridge for the subcutaneous administration of Troprelin Injection;â€—whether Mainbridge is able to complete the development of a new pump for the subcutaneous administration of Troprelin Injection and obtain FDA clearance on a timely basis or at all;â€—our cash resources;â€—the approvals or success of competitive products or technologies;â€—potential approvals of any product candidate we may develop, including YUTREPIA and L606, for marketing by the FDA or equivalent foreign regulatory authorities (and, if approved, the scope of the indications for which such product candidates are approved) or any failure to obtain such approvals;â€—our involvement in significant lawsuits, such as stockholder litigation, litigation involving the FDA, including the New FDA Litigation, or litigation related to intellectual property, including inter partes review proceedings and Hatch-Waxman litigation with originator companies or others which may hold patents, including the ongoing litigation in connection with the patents, trade secrets and confidential information that United Therapeutics has asserted against us;â€—regulatory or legal developments in the United States and other countries;â€—the results of our efforts to commercialize any product candidate we may develop, including YUTREPIA and L606, in the event we receive final approval from the FDA;â€—developments or disputes concerning patents or other proprietary rights;â€—the recruitment or departure of key personnel;â€—the level of expenses related to any of our product candidates or clinical development programs;â€—the results of our efforts to discover, develop, acquire or in-license additional product candidates or products;â€—actual or anticipated changes in estimates as to financial results, development timelines or recommendations by securities analysts;â€—variations in our financial results or those of companies that are perceived to be similar to us;â€—changes in the structure of healthcare payment systems;â€—market conditions in the pharmaceutical and biotechnology sectors and issuance of new or changed securities analystsâ€™ reports or recommendations;â€—general economic, industry and market conditions; and 75Table of Contentsâ€—the other factors described in this â€œRisk Factorsâ€ section.â€ The stock market in general, and market prices for the securities of pharmaceutical companies like ours in particular, have from time to time experienced volatility that often has been unrelated to the operating performance of the underlying companies. These broad market and industry fluctuations may adversely affect the market price of our Common Stock, regardless of our operating performance. Stock prices of many pharmaceutical companies have fluctuated in a manner unrelated or disproportionate to the operating performance of those companies. In several recent situations when the market price of a stock has been volatile, holders of that stock have instituted securities class action litigation against the company that issued the stock. If any of our stockholders were to bring a lawsuit against us, the defense and disposition of the lawsuit could be costly and divert the time and attention of our management and harm our operating results. Our principal stockholders and management own a significant percentage of our stock and will be able to exercise significant influence over matters subject to stockholder approval. Our executive officers, directors and principal stockholders, together with their respective affiliates, beneficially owned 39.0% of our capital stock as of October 30, 2024. Accordingly, our executive officers, directors and principal stockholders have significant influence in determining the composition of our board of directors (the â€œBoardâ€), and voting on all matters requiring stockholder approval, including mergers and other business combinations, and continue to have significant influence over our operations. This concentration of ownership could have the effect of delaying or preventing a change in our control or otherwise discouraging a potential acquirer from attempting to obtain control of us that you may believe are in your best interests as one of our stockholders. This in turn could have a material adverse effect on our stock price and may prevent attempts by our stockholders to replace or remove the Board or management. As a public company, we are obligated to develop and maintain proper and effective internal controls over financial reporting and any failure to do so may adversely affect investor confidence in us and, as a result, the trading price of our shares. Effective internal controls over financial reporting are necessary for us to provide reliable financial reports and, together with adequate disclosure controls and procedures, are designed to prevent fraud. Any failure to implement required new or improved controls, or difficulties encountered in their implementation could cause us to fail to meet our reporting obligations. Inferior internal controls could also cause investors to lose confidence in our reported financial information, which could have a negative effect on the trading price of our Common Stock. In addition, any future testing by us conducted in connection with Section 404 of the Sarbanes-Oxley Act of 2002, as amended (the â€œSarbanes-Oxley Actâ€) or the subsequent testing by our independent registered public accounting firm, may reveal deficiencies in our internal controls over financial reporting that are deemed to be material weaknesses or that may require prospective or retroactive changes to our consolidated financial statements or identify other areas for further attention or improvement.â€ As required by the Sarbanes Oxley Act and commencing with the fiscal year ended December 31, 2019, we were required to furnish a report by management on, among other things, the effectiveness of our internal controls over financial reporting. See Item 4. Controls and Procedures for additional information.â€ Because we are a â€œsmaller reporting company,â€ we may take advantage of certain scaled disclosures available to us, resulting in holders of our securities receiving less Company information than they would receive from a public company that is not a smaller reporting company. We are a â€œsmaller reporting companyâ€ as defined under Rule 12b-2 of the Exchange Act. As of December 31, 2023, we are no longer an â€œemerging growth company,â€ as defined in Section 2(a) of the Securities Act, as modified by the Jumpstart Our Business Startups Act of 2012. As a smaller reporting company, we may take advantage of certain of the scaled disclosures available to smaller reporting companies and will be able to take advantage of these scaled disclosures for so long as (i) our Common Stock held by non-affiliates is less than \$250 million measured on the last business day of our second fiscal quarter, or (ii) our annual revenue is less than \$100 million during the most recently completed fiscal year and our Common Stock held by non-affiliates is less than \$700 million measured on the last business day of our second fiscal quarter. Based on the closing price of our common stock on June 30, 2024 we will remain a smaller 76Table of Contents reporting company through at least the end of 2025. To the extent we take advantage of any reduced disclosure obligations, it may make it harder for investors to analyze the Companyâ€™s results of operations and financial prospectus in comparison with other public companies. As a smaller reporting company, we are permitted to comply with scaled-back disclosure obligations in our SEC filings compared to other issuers, including with respect to disclosure obligations regarding executive compensation in our periodic reports and proxy statements. We have elected to adopt the accommodations available to smaller reporting companies. Until we cease to be a smaller reporting company, the scaled-back disclosure in our SEC filings will result in less information about our company being available than for other public companies. If investors consider our Common Stock less attractive as a result of our election to use the scaled-back disclosure permitted for smaller reporting companies, there may be a less active trading market for our Common Stock and our share price may be more volatile. Anti-takeover provisions in our charter documents and under Delaware law could make an acquisition of us difficult, limit attempts by our stockholders to replace or remove our current management and adversely affect our stock price. Provisions of our certificate of incorporation and bylaws may delay or discourage transactions involving an actual or potential change in our control or change in our management, including transactions in which stockholders might otherwise receive a premium for their shares, or transactions that our stockholders might otherwise deem to be in their best interests. Therefore, these provisions could adversely affect the price of our stock. Among other things, the certificate of incorporation and bylaws:â€—permit the Board to issue up to 10 million shares of preferred stock, with any rights, preferences and privileges as they may designate;â€—provide that the authorized number of directors may be changed only by resolution of our Board;â€—provide that all vacancies, including newly created directorships, may, except as otherwise required by law, be filled by the affirmative vote of a majority of directors then in office, even if less than a quorum;â€—require that any action to be taken by our stockholders must be effected at a duly called annual or special meeting of stockholders and may not be taken by written consent;â€—create a staggered board of directors such that all members of our Board are not elected at one time;â€—allow for the issuance of authorized but unissued shares of our capital stock without any further vote or action by our stockholders; andâ€—establish advance notice requirements for nominations for election to the Board or for proposing matters that can be acted upon at stockholdersâ€™ meetings.â€ In addition, because we are incorporated in Delaware, we are governed by the provisions of Section 203 of the Delaware General Corporation Law (â€œDGCLâ€) which generally prohibits a Delaware corporation from engaging in any of a broad range of business combinations with any stockholder owning in excess of 15% of our outstanding stock for a period of three years following the date on which the stockholder obtained such 15% equity interest in us.â€ The terms of our authorized preferred stock selected by our Board at any point could decrease the amount of earnings and assets available for distribution to holders of our Common Stock or adversely affect the rights and powers, including voting rights, of holders of our Common Stock without any further vote or action by the stockholders. As a result, the rights of holders of our Common Stock will be subject to, and may be adversely affected by, the rights of the holders of any preferred stock that may be issued by us in the future, which could have the effect of decreasing the market price of our Common Stock.â€ Any provision of our certificate of incorporation or bylaws or Delaware corporate law that has the effect of delaying or deterring a change in control could limit opportunities for our stockholders to receive a premium for their shares of Common Stock, and could also affect the price that investors are willing to pay for our Common Stock.â€ 77Table of ContentsOur certificate of incorporation designates the Court of Chancery of the State of Delaware as the sole and exclusive forum for certain types of actions and proceedings that may be initiated by our stockholders, which could limit our stockholdersâ€™ ability to obtain a favorable judicial forum for disputes with us or our directors, officers or other employees. Our certificate of incorporation provides that, to the fullest extent permitted by law, the Court of Chancery of the State of Delaware will be the sole and exclusive forum for: (i) any derivative action or proceeding brought on our behalf; (ii) any action asserting a claim of breach of a fiduciary duty owed by any of our directors or officers to us or our stockholders; (iii) any action asserting a claim against us arising pursuant to any provision of the DGCL, our certificate of incorporation or our bylaws; or (iv) any action asserting a claim against us governed by the internal affairs doctrine; provided, that, this provision would not apply to suits brought to enforce a duty or liability created by the Securities Act or Exchange Act. Furthermore, our bylaws designate the federal district courts of the United States as the exclusive forum for the resolution of any complaint asserting a cause of action arising under the Securities Act. Any person or entity purchasing or otherwise acquiring any interest in shares of our capital stock is deemed to have received notice of and consented to the foregoing provisions. This choice of forum provision may limit a stockholderâ€™s ability to bring a claim in a judicial forum that it finds more favorable for disputes with us or our directors or officers, which may discourage such lawsuits against us and our directors or officers. Alternatively, if a court were to find this choice of forum provision inapplicable to, or unenforceable in respect of, one or more of the specified types of actions or proceedings, we may incur additional costs associated with resolving such matters in other jurisdictions, which could adversely affect our business, financial condition, prospects or results of operations.â€ Because we do not anticipate paying any cash dividends on our Common Stock in the foreseeable future, capital appreciation, if any, will be your sole source of gain. We have never declared or paid cash dividends on our equity securities. We currently intend to retain all of our future earnings, if any, to finance the growth and development of our business. In addition, the terms of our existing RIBA with HCR preclude us, and the terms of any future debt or financing agreement may preclude us, from paying dividends. As a result, capital appreciation, if any, of our equity securities will likely be your sole source of gain for the foreseeable future.â€ An impairment of our long-lived contract acquisition costs and intangible assets, including goodwill, could have a material non-cash adverse impact on our results of operations. In connection with the accounting for our RareGen acquisition, we have recorded significant amounts of contract acquisition costs, intangible assets, and goodwill. Under GAAP, we must assess, at least annually and potentially more frequently, whether the value of goodwill has been impaired. Contract acquisition costs and amortizing intangible assets will be assessed for impairment in the event of an impairment indicator. The valuation of goodwill depends on a variety of factors, the success of our business, including our ability to obtain regulatory approval for YUTREPIA, global market and economic conditions, earnings growth and expected cash flows. Impairments may be caused by factors outside our control, such as actions by the FDA, increasing competitive pricing pressures, and various other factors. Significant and unanticipated changes or our inability to obtain or maintain regulatory approvals for our product candidates, including the NDA for YUTREPIA, could require a non-cash charge for impairment in a future period, which may significantly affect our results of operations in the period of such charge.â€ General Risk FactorsGeneral Risks Related to the Commercialization of our Product Candidates Our business and operations may be adversely affected by the effects of health epidemics. Our business and operations could be adversely affected by health epidemics in regions where we have offices, manufacturing facilities, concentrations of clinical trial sites or other business operations, and could cause significant disruption in the operations of clinical trial sites, contract manufacturers or suppliers and contract research organizations upon whom we rely. 78Table of Contents The extent to which health epidemics impact our business and operations, including our clinical development and regulatory efforts, will depend on future developments that are highly uncertain and cannot be predicted with confidence at the time of this Quarterly Report on Form 10-Q, such as the severity and duration of future outbreaks, the duration and effect of business disruptions and the short-term effects, the administration, availability and efficacy of vaccination programs and the ultimate effectiveness of travel restrictions, quarantines, social distancing requirements and business closures in the United States and other countries to contain and treat any such health epidemic. These impacts could adversely affect our business, financial condition, results of operations and growth prospects. In addition, to the extent any health epidemic adversely affects our business and results of operations, it may also have the effect of heightening many of the other risks and uncertainties described in this â€œRisk Factorsâ€ section and the â€œRisk Factorsâ€ sections of the documents incorporated by reference herein. We are currently operating in a period of economic uncertainty and capital markets disruption, which has been significantly impacted by geopolitical instability. Changes and instability in global economic conditions and geopolitical matters could have a material adverse effect on our business, financial condition and results of operations. U.S. and global markets are experiencing volatility and disruption following the escalation of geopolitical tensions and the start of the military conflict between Russia and Ukraine. In February 2022, a full-scale military invasion of Ukraine by Russian troops began. Although the length and impact of the ongoing military conflict is highly unpredictable, the conflict in Ukraine has led to market disruptions, including significant volatility in commodity prices, credit and capital markets, as well as supply chain interruptions, which has contributed to periods of high inflation globally. We are continuing to monitor the situation in

Ukraine and global capital markets and assessing its potential impact on our business. The global economy has been, and may continue to be, negatively impacted by Russia's invasion of Ukraine. As a result of Russia's invasion of Ukraine, the U.S., the European Union, the United Kingdom, and other G7 countries, among other countries, have imposed substantial financial and economic sanctions on certain industry sectors and parties in Russia. Broad restrictions on exports to Russia have also been imposed. These measures include: (i) comprehensive financial sanctions against major Russian banks; (ii) additional designations of Russian individuals with significant business interests and government connections; (iii) designations of individuals and entities involved in Russian military activities; and (iv) enhanced export controls and trade sanctions limiting Russia's ability to import various goods. Russian military actions and the resulting sanctions could continue to adversely affect the global economy and financial markets and lead to instability and lack of liquidity in capital markets, potentially making it more difficult for us to obtain additional funds. In addition, on October 7, 2023, Hamas militants and members of other terrorist organizations infiltrated Israel's southern border from the Gaza Strip and conducted a series of terror attacks on civilian and military targets. Thereafter, Hamas launched extensive rocket attacks on Israeli population and industrial centers located along the Israeli border with the Gaza Strip. Shortly following the attack, Israel's security cabinet declared war against Hamas and launched an aerial bombardment of various targets within the Gaza Strip. The Israeli government subsequently called for the evacuation of over one million residents of the northern part of the Gaza Strip and initiated ground operations in the Gaza Strip. It is possible that other terrorist and/or regional organizations will join the hostilities as well, including Hezbollah in Lebanon, and Palestinian military organizations in the West Bank, resulting in a widening of the conflict. The intensity and duration of Israel's current war against Hamas is difficult to predict as are such war's economic implications on the global economy. In addition, since the commencement of these events, there have been continued hostilities along Israel's northern border with Lebanon (with the Hezbollah terror organization) and southern border (with the Houthi movement in Yemen). It is possible that hostilities with Hezbollah in Lebanon will escalate, and that other terrorist organizations, including Palestinian military organizations in the West Bank as well as other hostile countries will join the hostilities. Such clashes may escalate in the future into a greater regional conflict.

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Furthermore, because of current geopolitical tensions, the Biden administration has recently signed multiple executive orders regarding China. One particular executive order titled Advancing Biotechnology and Biomanufacturing Innovation for a Sustainable, Safe, and Secure American Bioeconomy, signed on September 12, 2022, will likely impact the pharmaceutical industry to encourage U.S. domestic manufacturing of pharmaceutical products. Moreover, there have been Congressional legislative proposals, such as the recent bill titled the BIOSECURE Act, to discourage contracting with Chinese companies on the development or manufacturing of pharmaceutical products. The BIOSECURE Act passed the U.S. House of Representatives on September 9, 2024. The version of the BIOSECURE Act that passed the U.S. House of Representatives included a grandfather clause that would allow contracts entered into with the Chinese companies named therein prior to the effective date of such legislation until January 1, 2032. The BIOSECURE Act must also pass the U.S. Senate before going to President Biden for either his veto or signature, and it is uncertain whether the bill will be brought to the floor for a vote by the U.S. Senate before the current legislative session expires on January 3, 2025. Any additional executive orders or legislative action regarding or potential sanctions on China could materially impact our current manufacturing partners. Although our business has not been materially impacted by these geopolitical tensions to date, such matters may affect our business and it is impossible to predict the extent to which our operations, or those of our suppliers and manufacturers, will be impacted in the short and long term, or the ways in which such matters may impact our business. The extent and duration of the military action, sanctions, actual or perceived political instability and resulting market disruptions are impossible to predict but could be substantial. Any such disruptions may also magnify the impact of other risks described herein. The U.S. political and economic environment could materially impact our business operations and financial performance, and uncertainty surrounding the potential legal, regulatory and policy changes by a new U.S. presidential administration may directly affect us and the global economy. The political and economic environment in the U.S. and elsewhere has resulted in and will continue to result in some uncertainty. Changing regulatory policies because of the changing political environment could impact our regulatory and compliance costs and future revenues, all of which could materially and adversely affect our business, financial condition and operating results. Failure to adapt to or comply with evolving regulatory requirements or investor or stakeholder expectations and standards could negatively impact our reputation, ability to do business with certain partners, access to capital and our stock price. Further, the recent presidential election and congressional seat turnover may result in increased regulatory and economic uncertainty. Changes in federal policy by the executive branch and regulatory agencies may occur over time through the new presidential administration's and/or Congress's policy and personnel changes, which could lead to changes involving the level of oversight and focus on the pharmaceutical industry; however, the nature, timing and economic and political effects of such potential changes remain highly uncertain. Any future changes in federal and state laws and regulations, as well as the interpretation and implementation of such laws and regulations, could affect us in substantial and unpredictable ways. At this time, it is unclear what laws, regulations and policies may change and whether future changes or uncertainty surrounding future changes will adversely affect our operating environment and therefore our business, financial condition and results of operations. If the FDA or comparable regulatory authorities in other countries approve generic versions of our product candidates, or do not grant our product candidates a sufficient period of market exclusivity before approving their generic versions, our ability to generate revenue may be adversely affected. Once an NDA is approved, the drug product covered will be listed as a reference listed drug in the FDA's Orange Book. In the United States, manufacturers of drug products may seek approval of generic versions of reference listed drugs through the submission of abbreviated new drug applications, or ANDAs. In support of an ANDA, a generic manufacturer is generally required to show that its product has the same active pharmaceutical ingredient(s), dosage form, strength, route of administration and conditions of use or labeling as the reference listed drug and that the generic version is bioequivalent to the reference listed drug. Generic drug products may be significantly less expensive to bring to market than the reference listed drug, and companies that produce generic drug products are generally able to offer 80Table of Contents them at lower prices. Thus, following the introduction of a generic drug product, a significant percentage of the sales of any reference listed drug may be lost to the generic drug product. The FDA will not approve an ANDA for a generic drug product until the applicable period of market exclusivity for the reference listed drug has expired. The applicable period of market exclusivity varies depending on the type of exclusivity granted. A grant of market exclusivity is separate from the existence of patent protection and manufacturers may seek to launch generic versions of our drug products following the expiry of their respective marketing exclusivity periods, even if our drug products are still under patent protection at the relevant time. Any competition that our product candidates may face, if and when such product candidates are approved for marketing and commercialized, from generic versions could substantially limit our ability to realize a return on our investment in the development of our product candidates and have a material and adverse effect on our business and prospects. We are subject to risks related to information technology systems, including cyber-security risks; successful cyber-attacks or technological malfunctions can result in, among other things, financial losses, the inability to process transactions, the unauthorized release of confidential information and reputational risk, all of which would negatively impact our business, financial condition or results of operations. Our use of technology is critical to our continued operations. We are susceptible to operational, financial and information security risks resulting from cyber-attacks or technological malfunctions. Successful cyber-attacks or technological malfunctions affecting us, our CMOS or our business partners can result in, among other things, financial losses, the inability to process transactions, the unauthorized release of confidential or proprietary information and reputational risk. As cybersecurity threats continue to evolve, we may be required to use additional resources to continue to modify or enhance protective measures or to investigate security vulnerabilities, which could have a material adverse effect on our business, financial condition or results of operations.

General Risks Related to the Development and Regulatory Approval of our Product Candidates

Even if we obtain marketing approval for our product candidates in the United States, we or our collaborators may not obtain marketing approval for the same product candidates elsewhere. We may enter into strategic collaboration arrangements with third parties to commercialize our product candidates outside of the United States. In order to market any product candidate outside of the United States, we or our collaborators will be required to comply with numerous and varying regulatory requirements of other countries regarding safety and efficacy. Clinical trials conducted in one country may not be recognized or accepted by regulatory authorities in other countries, and obtaining marketing approval in one country does not mean that marketing approval will be obtained in any other country. Approval processes vary among countries and additional product testing and validation, or additional administrative review periods, may be required from one country to the next. Seeking marketing approval in countries other than the United States could be costly and time-consuming, especially if additional preclinical studies or clinical trials are required to be conducted. We currently do not have any product candidates approved for sale in any jurisdiction, including non-U.S. markets, and we do not have experience in obtaining marketing approval in non-U.S. markets. We currently also have not identified any collaborators to market our products outside of the United States and cannot assure you that such collaborators, even if identified, will be able to successfully obtain marketing approval for our product candidates outside of the United States. If we or our collaborators fail to obtain marketing approval in non-U.S. markets, or if such approval is delayed, our target market may be reduced, and our ability to realize the full market potential of our products will be adversely affected.

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General Risks Related to Healthcare Regulation

The pharmaceutical industry is subject to a range of laws and regulations in areas including healthcare program requirements and fraud, waste, and abuse; healthcare and related marketing compliance and transparency; and privacy and data security. Our failure to comply with these laws and regulations as they are, or in the future become, applicable to us may have an adverse effect on our business. Healthcare providers, physicians and third-party payors often play a primary role in the recommendation and prescription of any drug products for which we may obtain marketing approval, or for which we may provide contracted promotional services to third parties. Our current and future arrangements with healthcare providers, physicians, third-party payors and customers, and our sales, marketing and educational activities, may expose us to broadly applicable fraud and abuse and other healthcare laws and regulations (at the federal and state level) that may constrain our business or financial arrangements and relationships through which we market, sell, or distribute drug products.

If in addition, we may be subject to transparency laws and patient privacy regulation by both the federal government and the states in which we conduct our business. We also plan to conduct clinical trials and may in the future conduct business in jurisdictions outside of the United States, which may cause us to become subject to transparency law and privacy regulations in those jurisdictions as well.

The laws that may affect our ability to operate include, but are not limited to, the following examples:

- The federal Anti-Kickback Statute, or AKS, prohibits, among other things, persons and entities including pharmaceutical manufacturers from, among other things, knowingly and willfully soliciting, receiving, offering or paying remuneration, directly or indirectly, overtly or covertly, in cash or in kind, to induce or reward, or in return for, either the referral of an individual for or the purchase, lease, or order of, or the arranging for an item or service for which payment may be made, in whole or in part, under federal healthcare programs such as the Medicare and Medicaid programs.
- The federal civil and criminal false claims laws and civil monetary penalty laws impose a range of prohibitions and compliance considerations. For example, the False Claims Act, or the FCA, prohibits individuals or entities from, among other things, knowingly presenting, or causing to be presented, claims for payment to, or approval by, the federal government that are false, fictitious or fraudulent or knowingly making, using or causing to be made or used, a false record or statement material to a false or fraudulent claim to avoid, decrease or conceal an obligation to pay money to the federal government. Claims resulting from a violation of the federal AKS constitute a false or fraudulent claim for purposes of the FCA. Promotion that is deemed to be a false or fraudulent claim can be the basis of FCA exposure.
- Federal law includes provisions (established under the Health Insurance Portability and Accountability Act of 1996) addressing healthcare fraud and false statements relating to healthcare matters. The healthcare fraud statute prohibits knowingly and willfully executing, or attempting to execute, a scheme to defraud any healthcare benefit program, including private payors. The false statements statute prohibits knowingly and willfully falsifying, concealing or covering up a material fact or making any materially false, fictitious or fraudulent statement in connection with the delivery of or payment for healthcare benefits, items or services. Violations of these statutes is a felony and may result in fines, imprisonment or exclusion from governmental programs.
- Privacy and data security laws may apply to our business. Under Section 5(a) of the Federal Trade Commission Act, the Federal Trade Commission expects a company's data security measures to be reasonable and appropriate in light of the sensitivity and volume of consumer information it holds, the size and complexity of its business, and the cost of available tools to improve security and reduce vulnerabilities. Medical data is considered sensitive data that merits stronger safeguards. States may also impose requirements, for example the California Consumer Privacy Act created data privacy obligations for covered companies and providing privacy rights to California residents, including the right to opt out of certain disclosures of their information. In addition, if we engage in business activities outside of the United States, including clinical trials that we plan to conduct outside of the United States, we may become subject to privacy and data security laws in those additional jurisdictions in which we operate or conduct clinical trials.
- The federal physician payment transparency requirements, sometimes referred to as the Physician Payments Sunshine Act, requires applicable manufacturers of covered drugs, devices, biologics and medical supplies for which payment is available under government healthcare programs to annually report to the Centers for Medicare and Medicaid Services, or the CMS, information related to certain payments or other transfers of value made or distributed to physicians and teaching hospitals, as well as ownership and investment interests held by physicians and their immediate family members. Payments and transfers of value made to certain other providers such as nurse practitioners and physician assistants will also need to be reported under the Sunshine Act.
- For both investigational and commercialized products, interactions with or communications directed to healthcare professionals, patients or patient- or disease-advocates or advocacy groups, and payors, are subject to heightened scrutiny by the FDA. Relative to non-promotional communications, for example, there are specific and limited FDA accommodations for non-promotional, truthful and non-misleading sharing of information regarding products in development and off-label uses including dissemination of peer-reviewed reprints, support of independent continuing medical education, and healthcare economic discussions with payors. In a competitive environment, a company's communications about products in development may also be subject to heightened scrutiny.
- Analogous state laws and regulations, such as state anti-kickback and false claims laws, may apply to items or services reimbursed by any third-party payor, including commercial insurers, and in some cases may apply regardless of payor (i.e., even for self-pay scenarios). Some state laws require pharmaceutical companies to comply with the pharmaceutical industry's voluntary compliance guidelines and the relevant compliance guidance promulgated by the federal government in addition to requiring drug manufacturers to report pricing and marketing information, including, among other things, information related to payments to physicians and other healthcare providers or marketing expenditures, state and local laws that require the registration of pharmaceutical sales representatives. Many of these state laws differ from each other in significant ways and may not have the same effect, and may apply more broadly or be stricter than their federal counterparts, thus complicating compliance efforts; and
- Price reporting laws require the calculation and reporting of complex pricing metrics to government programs, where such reported prices may be used in the calculation of reimbursements or discounts on our drug products. Participation in such programs and compliance with their requirements may subject us to increased infrastructure costs and potentially limit our ability to price our drug products.
- Ensuring that our business and business arrangements with third parties comply with applicable healthcare laws, as well as responding to possible investigations by government authorities, can be time- and resource-consuming and can divert management's attention from the business, even if the government ultimately finds that no violation has occurred. If our operations are found to be in violation of any of the laws or regulations described above or any other laws or government regulations that apply to us, we may be subject to penalties and potentially, the curtailment or restructuring of our operations as well as additional governmental reporting obligations and oversight, any of which could adversely affect our ability to operate our business and our results of operations.

Recently enacted and future legislation and other legal developments may increase the difficulty and cost for us to obtain marketing approval of and commercialize our products and product candidates and affect the prices we may obtain.

In the United States and some foreign jurisdictions, there have been a number of legislative and regulatory changes and proposed changes regarding the healthcare system that could prevent or delay marketing approval of our product candidates.

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candidates, restrict or regulate post-approval activities and affect our ability to profitably sell any product candidates for which we obtain marketing approval.

In the United States, the Patient Protection and Affordable Care Act, as amended by the Health Care and Education Affordability Reconciliation Act, or collectively the ACA, is a sweeping law intended to broaden access to health insurance, reduce or constrain the growth of healthcare spending, enhance remedies against fraud and abuse, add new transparency requirements for the healthcare and health insurance industries, impose taxes and fees on the health industry and impose additional health policy reforms.

Among the provisions of the ACA of importance to our product candidates are the following:

- establishment of a new pathway for approval of lower-cost biosimilars to compete with biologic products;
- an increase in the statutory minimum rebates a manufacturer must pay under the Medicaid Drug Rebate Program;
- a new Medicare Part D coverage gap discount program, in which manufacturers must agree to offer point-of-sale discounts

off negotiated prices;—extension of manufacturers' Medicaid rebate liability;—expansion of eligibility criteria for Medicaid programs;—expansion of the entities eligible for discounts under the Public Health Service pharmaceutical pricing program;—a new requirement to annually report drug samples that manufacturers and distributors provide to physicians; and—**a new Patient-Centered Outcomes Research Institute to oversee, identify priorities in and conduct comparative clinical effectiveness research, along with funding for such research.** Since its enactment, there have been judicial, executive and Congressional challenges to certain aspects of the ACA. On June 17, 2021, the U.S. Supreme Court dismissed the most recent judicial challenge to the ACA brought by several states without specifically ruling on the constitutionality of the ACA. Prior to the Supreme Court's decision, President Biden issued an executive order initiating a special enrollment period from February 15, 2021 through August 15, 2021 for purposes of obtaining health insurance coverage through the ACA marketplace. The executive order also instructed certain governmental agencies to review and reconsider their existing policies and rules that limit access to healthcare. In addition, other legislative changes have been proposed and adopted since the ACA was enacted. For example, the Budget Control Act of 2011, enacted in August 2011, required sequestration that included aggregate reductions of Medicare payments to providers, which went into effect on April 1, 2013 and, due to subsequent legislative amendments, will remain in effect through 2032, unless additional Congressional action is taken. Under current legislation, the actual reduction in Medicare payments will increase in future years of the sequester. On January 2, 2013, the American Taxpayer Relief Act of 2012 was signed into law, which, among other things, reduced Medicare payments to several providers, including hospitals, and an increase in the statute of limitations period for the government to recover overpayments to providers from three to five years. Further, in March 2021, the American Rescue Plan Act of 2021 was signed into law, which, among other things, eliminated the statutory cap on drug manufacturers' Medicaid Drug Rebate Program rebate liability, effective January 1, 2024. Under current law enacted as part of the ACA, drug manufacturers' Medicaid Drug Rebate Program rebate liability is capped at 100% of the average manufacturer price for a covered outpatient drug. In addition, on September 20, 2024, the Centers for Medicare & Medicaid Services issued a final rule titled "Medicaid Program: Misclassification of Drugs, Program Integrity Updates Under the Medicaid Drug Rebate Program" which may impact our reimbursement and rebate strategy. We expect that other healthcare reform measures that may be adopted in the future may result in additional reductions in Medicare and other healthcare funding, more rigorous coverage criteria, new payment methodologies and in additional downward pressure on the price that we receive for any approved product. Any reduction in reimbursement from Medicare or other government programs may result in a similar reduction in payments from private payors. The implementation of cost containment measures or other healthcare reforms may prevent us from 84Table of Contentsbeing able to price our products at what we consider to be a fair or competitive price, generate revenue, attain profitability, or commercialize our product candidates, if approved. Moreover, there has recently been heightened governmental scrutiny over the manner in which manufacturers set prices for their marketed products. Individual states in the United States have become increasingly active in implementing regulations designed to contain pharmaceutical and biological product pricing, including price or patient reimbursement constraints, discounts, restrictions on certain product access and marketing cost disclosure and transparency measures. Most significantly, on August 16, 2022, President Biden signed the Inflation Reduction Act of 2022, or the IRA, into law. This statute marks the most significant action by Congress with respect to the pharmaceutical industry since adoption of the ACA in 2010. Among other things, the IRA requires manufacturers of certain drugs to engage in price negotiations with Medicare (beginning in 2026), with prices that can be negotiated subject to a cap; imposes rebates under Medicare Part B and Medicare Part D to penalize price increases that outpace inflation; and replaces the Part D coverage gap discount program with a new discounting program (beginning in 2025). The IRA permits the Secretary of the Department of Health and Human Services, or HHS, to implement many of these provisions through guidance, as opposed to regulation, for the initial years. HHS has and will continue to issue and update guidance as these programs are implemented. On August 29, 2023, HHS announced the list of the first ten drugs that will be subject to price negotiations, although the Medicare drug price negotiation program is currently subject to legal challenges. Further, the Biden administration released an additional executive order on October 14, 2022, directing HHS to submit a report within 90 days on how the Center for Medicare and Medicaid Innovation can be further leveraged to test new models for lowering drug costs for Medicare and Medicaid beneficiaries. In response to the executive order, on February 14, 2023, HHS released a report outlining three new models for testing by the CMS Innovation Center which will be evaluated on their ability to lower the cost of drugs, promote accessibility and improve quality of care. It is unclear whether the models will be utilized in any health reform measures in the future. Legally mandated price controls on payment amounts by third-party payors or other restrictions could harm our ability to price our products appropriately, which could negatively impact our business, results of operations, financial condition and prospects. In addition, regional healthcare authorities and individual hospitals are increasingly using bidding procedures to determine what pharmaceutical products and which suppliers will be included in their prescription drug and other healthcare programs. This could reduce the ultimate demand for our product candidates, if approved, or put pressure on our product pricing, which could negatively affect our business, results of operations, financial condition and prospects. Legislative and regulatory proposals have been made to expand post-approval requirements and restrict sales and promotional activities for pharmaceutical products. We cannot be sure whether additional legislative changes will be enacted, or whether the FDA or foreign regulations, guidance or interpretations will be changed, or what the impact of such changes on the marketing approvals of our product candidates, if any, may be. In addition, increased scrutiny by Congress of the FDA's approval process may significantly delay or prevent marketing approval, as well as subject us to more stringent product labeling and post-marketing testing and other requirements. There is also a great degree of uncertainty regarding how the recent U.S. Supreme Court decisions, including *Loper Bright Enterprises v. Raimondo and Corner Post, Inc. v. Board of Governors of the Federal Reserve System*, will impact FDA's enforcement and decision-making authority. Loper Bright explicitly overturned Chevron deference, which previously gave judicial deference to administrative action by agencies in the executive branch. Further, the Supreme Court's decision in Corner Post may result in challenges to FDA decisions by new litigants long into the future, resulting in greater uncertainty about our continued operations. **General Risks Related to Our Dependence on Third Parties** We rely on third parties to conduct our preclinical studies and clinical trials. We currently rely on, and plan to continue to rely on, third-party contract research organizations, or CROs, to monitor and manage data for our preclinical studies and clinical trials. However, we are responsible for ensuring that each of our trials is conducted in accordance with the applicable regulatory standards and our reliance on CROs does not relieve us of our regulatory responsibilities. The CROs on which we rely are required to comply with FDA regulations (and the regulations of comparable regulatory authorities in other countries) regarding GCP. Regulatory authorities enforce GCP standards through periodic 85Table of Contentsinspections. If any of the CROs on which we rely fail to comply with the applicable GCP standards, the clinical data generated in our clinical trials may be deemed unreliable. While we have contractual agreements with these CROs, we have limited influence over their actual performance and cannot control whether or not they devote sufficient time and resources to our preclinical studies and clinical trials. A failure to comply with the applicable regulations in the conduct of the preclinical studies and clinical trials for our product candidates may require us to repeat such studies or trials, which would delay the process of obtaining marketing approval for our product candidates and have a material and adverse effect on our business and prospects. Some of our CROs have the ability to terminate their respective agreements with us if, among others, it can be reasonably demonstrated that the safety of the patients participating in our clinical trials warrants such termination. If any of our agreements with our CROs is terminated, and if we are not able to enter into agreements with alternative CROs on acceptable terms or in a timely manner, or at all, the clinical development of our product candidates may be delayed and our development expenses could be increased. **General Risks Related to Legal Compliance Matters** Even if we obtain regulatory approval for a product candidate, our products and business will remain subject to ongoing regulatory obligations and review. If our product candidates are approved, they will be subject to ongoing regulatory requirements for manufacturing, labeling, packaging, storage, drug supply chain security surveillance and tracking, advertising, promotion, sampling, record-keeping, conduct of post-marketing studies and submission of safety, efficacy and other post-market information, including both federal and state requirements in the United States and comparable requirements outside of the United States. Accordingly, we and others with whom we work must continue to expend time, money and effort in all areas of regulatory compliance, including manufacturing, production and quality control. Any regulatory approvals that we may receive for our product candidates may also be subject to limitations on the approved indicated uses for which the product may be marketed or to the conditions of approval, or contain requirements for potentially costly post-marketing testing, including Phase 4 clinical trials, and surveillance to monitor the safety and efficacy of the product candidate. The FDA may also require a REMS as a condition of approval of our product candidates, which could include requirements for a medication guide, physician communication plans or additional elements to ensure safe use, such as restricted distribution methods, patient registries and other risk minimization tools. We will also be required to report certain adverse reactions and production problems, if any, to the FDA or other regulatory agencies and to comply with requirements concerning advertising and promotion for our products. Promotional communications with respect to prescription drugs are subject to a variety of legal and regulatory restrictions and must be consistent with the information in the product's approved label. As such, we may not promote our products for indications or uses for which they do not have FDA or other regulatory agency approval. The holder of an approved NDA must also submit new or supplemental applications and obtain FDA approval for certain changes to the approved product, product labeling, or manufacturing process. We could also be asked to conduct post-marketing clinical studies to verify the safety and efficacy of our product candidates in general or in specific patient subsets. An unsuccessful post-marketing study or failure to complete such a clinical study could result in the withdrawal of marketing approval. Furthermore, any new legislation addressing drug safety issues could result in delays in product development or commercialization or increased costs to assure compliance. Foreign regulatory authorities impose similar requirements. If a regulatory agency discovers previously unknown problems with a product, such as adverse events of unanticipated severity or frequency, or disagrees with the promotion, marketing or labeling of a product, such regulatory agency may impose restrictions on that product or us, including requiring withdrawal of the product from the market. If we fail to comply with applicable regulatory requirements, regulatory agency or enforcement authority may, among other things:—issue warning letters asserting that we are in violation of the law;—seek an injunction or impose civil or criminal penalties or monetary fines;—suspend or withdraw regulatory approval;—suspend any of our ongoing clinical trials;—refuse to approve pending applications or supplements to approved applications submitted by us or our strategic partners;—restrict the marketing or manufacturing of our products; 86Table of Contents—seize or detain products, or require a product recall;—refuse to permit the import or export of our product candidates; or—refuse to allow us to enter into government contracts. Any government investigation of alleged violations of law could require us to expend significant time and resources in response and could generate negative publicity. Any failure to comply with ongoing regulatory requirements may significantly and adversely affect our ability to commercialize and generate revenue from our product candidates. If regulatory sanctions are applied or if regulatory approval is withdrawn, the value of our company and our operating results will be adversely affected. We also cannot predict the likelihood, nature or extent of government regulation that may arise from future legislation or administrative or executive action, either in the United States or abroad. If we are slow or unable to adapt to changes in existing requirements or the adoption of new requirements or policies, or if we are not able to maintain regulatory compliance, we may lose any marketing approval that we may have obtained and we may not achieve or sustain profitability, which would adversely affect our business, prospects, financial condition and results of operations. Environmental, social and governance matters may impact our business and reputation. Governmental authorities, non-governmental organizations, customers, investors, external stakeholders and employees are increasingly sensitive to environmental, social and governance, or ESG, concerns, such as diversity and inclusion, climate change, water use, recyclability or recoverability of packaging, and plastic waste. This focus on ESG concerns may lead to new requirements that could result in increased costs associated with developing, manufacturing and distributing our products. Our ability to compete could also be affected by changing customer preferences and requirements, such as growing demand for more environmentally friendly products, packaging or supplier practices, or by failure to meet such customer expectations or demand. While we strive to improve our ESG performance, we risk negative stockholder reaction, including from proxy advisory services, as well as damage to our brand and reputation, if we do not act responsibly, or if we are perceived to not be acting responsibly in key ESG areas, including equitable access to medicines and vaccines, product quality and safety, diversity and inclusion, environmental stewardship, support for local communities, corporate governance and transparency, and addressing human capital factors in our operations. If we do not meet the ESG expectations of our investors, customers and other stakeholders, we could experience reduced demand for our products, loss of customers, and other negative impacts on our business and results of operations. Climate change or legal, regulatory or market measures to address climate change may negatively affect our business, results of operations, cash flows and prospects. We believe that climate change has the potential to negatively affect our business and results of operations, cash flows and prospects. We are exposed to physical risks (such as extreme weather conditions or rising sea levels), risks in transitioning to a low-carbon economy (such as additional legal or regulatory requirements, changes in technology, market risk and reputational risk) and social and human effects (such as population dislocations and harm to health and well-being) associated with climate change. These risks can be either acute (short-term) or chronic (long-term). The adverse impacts of climate change include increased frequency and severity of natural disasters and extreme weather events such as hurricanes, tornados, wildfires (exacerbated by drought), flooding, and extreme heat. Extreme weather and sea-level rise pose physical risks to our facilities as well as those of our suppliers. Such risks include losses incurred as a result of physical damage to facilities, loss or spoilage of inventory, and business interruption caused by such natural disasters and extreme weather events. Other potential physical impacts due to climate change include reduced access to high-quality water in certain regions and the loss of biodiversity, which could impact future product development. These risks could disrupt our operations and its supply chain, which may result in increased costs. New legal or regulatory requirements may be enacted to prevent, mitigate, or adapt to the implications of a changing climate and its effects on the environment. These regulations, which may differ across jurisdictions, could result in us being subject to new or expanded carbon pricing or taxes, increased compliance costs, restrictions on greenhouse gas emissions, investment in new technologies, increased carbon disclosure and transparency, upgrade of facilities to meet 87Table of Contentsnew building codes, and the redesign of utility systems, which could increase our operating costs, including the cost of electricity and energy used by us. Our supply chain would likely be subject to these same transitional risks and would likely pass along any increased costs to us. **General Risks Related to our Intellectual Property** We may become involved in litigation to protect our intellectual property or enforce our intellectual property rights, which could be expensive, time-consuming and may not be successful. Competitors may infringe our patents or misappropriate or otherwise violate our intellectual property rights. To counter infringement or unauthorized use, we may engage in litigation to, among others, enforce or defend our intellectual property rights, determine the validity or scope of our intellectual property rights and those of third parties, and protect our trade secrets. Such actions may be time-consuming and costly and may divert our management's attention from our core business and reduce the resources available for our clinical development, manufacturing and marketing activities, and consequently have a material and adverse effect on our business and prospects, regardless of the outcome. In addition, in an infringement proceeding, a court may decide that a patent owned by, or licensed to, us is invalid or unenforceable, or may refuse to stop the other party from using the technology in question on the ground that our patents do not cover such technology. An adverse result in any litigation proceeding could put one or more of our patents at risk of being invalidated, held unenforceable or interpreted narrowly. Furthermore, because of the substantial amount of discovery required in connection with intellectual property litigation, there is a risk that our confidential information may be compromised by disclosure. Patent terms may be inadequate to protect our competitive position on our product candidates for an adequate amount of time. Patents have a limited lifespan. In the United States, the natural expiration of a patent is generally 20 years after it is filed. While various extensions may be available, the life of a patent, and the protection it affords, is limited. Given the amount of time required for the development, testing and regulatory review of new product candidates, patents protecting such candidates might expire before or shortly after such candidates are commercialized. We intend to seek extensions of patent terms in the United States and, if available, in other countries where we prosecute patents. In the United States, the Hatch-Waxman Act permits patent owners to request a patent term extension, based on the regulatory review period for a product, of up to five years beyond the normal expiration of the patent, which is limited to one patent claiming the approved drug product or use in an indication (or any additional indications approved during the period of extension). However, the applicable authorities, including the FDA and the USPTO, in the United States, and comparable regulatory authorities in other countries, may not agree with our assessment of whether such extensions are available, and may refuse to grant extensions to our patents, or grant more limited extensions than we had requested. In such event, our competitors may be able to take advantage of our investment in development and clinical trials by referencing our preclinical and clinical data in their marketing approval applications with the FDA to launch their drug product earlier than might otherwise be the case. **General Risks Related to the Manufacturing of our Product Candidates** Our facilities are subject to extensive and ongoing regulatory requirements and failure to comply with these regulations may result in significant liability. Our company and our facilities are subject to payment of fees, registration and listing requirements, ongoing review and periodic inspections by the FDA and other regulatory

authorities for compliance with quality system regulations, including the FDA's cGMP requirements. These regulations cover all aspects of the manufacturing, testing, quality control and record-keeping of our drug products. Furthermore, the facilities where our product candidates are manufactured may be subject to additional inspections by the FDA before we can obtain final marketing approval and remain subject to periodic inspection even after our product candidates have received marketing approval. Suppliers of 88Table of Contentscomponents and materials, such as active pharmaceutical ingredients, used to manufacture our drug products are also required to comply with the applicable regulatory standards. The manufacture of pharmaceutical products is complex and requires significant expertise and capital investment, including the development of advanced manufacturing techniques and process controls. We and any contract manufacturers that we may engage in the future must comply with cGMP requirements. Manufacturers of pharmaceutical products often encounter difficulties in production, particularly in scaling up and validating initial production and contamination controls. These problems include difficulties with production costs and yields, quality control, including stability of the product, quality assurance testing, operator error, shortages of qualified personnel, as well as compliance with strictly enforced federal, state and foreign regulations. Furthermore, if microbial, viral or other contaminations are discovered in our product candidates or in the manufacturing facilities in which our product candidates are made, such manufacturing facilities may need to be closed for an extended period of time to investigate and remedy the contamination. Compliance with these regulatory standards often requires significant expense and effort. If we or our suppliers are unable to comply with the applicable regulatory standards or take satisfactory corrective steps in response to adverse results of an inspection, this could result in enforcement action, including, among others, the issue of a public warning letter, a shutdown or of restrictions on our or our suppliers' manufacturing operations, delays in approving our drug products and refusal to permit the import or export of our drug products. Any adverse regulatory action taken against us could subject us to significant liability and harm our business and prospects. Item 5. Other Information Rule 10b5-1 Trading Plans During the three months ended September 30, 2024, none of our directors or officers (as defined in Rule 16a-1(f) under the Exchange Act) adopted or terminated any contract, instruction or written plan for the purchase or sale of our securities that was intended to satisfy the affirmative defense conditions of Rule 10b5-1(c) under the Exchange Act or any non-Rule 10b5-1 trading arrangement as defined in Item 408(c) of Regulation S-K. During the three months ended September 30, 2024, the Company did not adopt or terminate a Rule 10b5-1 trading arrangement (as defined in Item 408(a)(1)(i) of Regulation S-K). Item 6. Exhibits The exhibits listed on the Exhibit A Index hereto are filed or furnished (as stated therein) as part of this Quarterly Report on Form 10-Q. Exhibit No. A-1 is a Document 10.1*^a Common Stock Purchase Agreement by and among Liquida Corporation and the Purchasers, dated September 10, 2024 (incorporated by reference to Exhibit 10.1 of the Company's Current Report on Form 8-K, filed with the SEC on September 12, 2024). Item 10.2*^a Registration Rights Agreement by and among Liquida Corporation and the Purchasers, dated September 10, 2024 (incorporated by reference to Exhibit 10.2 of the Company's Current Report on Form 8-K, filed with the SEC on September 12, 2024). Item 10.3*^a Fifth Amendment to Revenue Interest Financing Agreement, dated as of September 11, 2024, by and between Liquida Technologies, Inc. and Healthcare Royalty Partners IV, L.P. (incorporated by reference to Exhibit 10.3 of the Company's Current Report on Form 8-K, filed with the SEC on September 12, 2024). Item 10.4*^a Device License Agreement, dated as of October 2, 2024, by and between Liquida Technologies, Inc. and Pharmosa Biopharm Inc. Item 10.5*^a First Amendment to the License Agreement, dated as of October 2, 2024, by and between Liquida Technologies, Inc. and Pharmosa Biopharm Inc. Item 31.1*^a Certification of Principal Executive Officer pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 302 of the Sarbanes-Oxley Act. Item 31.2*^a Certification of Principal Financial Officer pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act. Item 32.2*^a Certification of Principal Financial Officer pursuant to 18 U.S.C. Section 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act. Item 101.INS*^a Inline XBRL Instance Document 101.SCH*^a Inline XBRL Taxonomy Extension Schema Document 101.CAL*^a Inline XBRL Taxonomy Extension Calculation Linkbase Document 101.DEF*^a Inline XBRL Taxonomy Extension Definition Linkbase Document 101.LAB*^a Inline XBRL Taxonomy Extension Label Linkbase Document 104*^a Cover Page A Interactive Data File (formatted in Inline XBRL and contained in Exhibit A 101). Filed herewith. **Furnished herewith. ++Portions of this exhibit have been redacted in compliance with Regulation S-K Item 601(b)(10). The omitted information is not material and would likely cause competitive harm to the Company if publicly disclosed. Item 90 Table of Contents SIGNATURES Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned duly authorized. DATE: November 13, 2024. By/s/ Roger A. Jeffs, Ph.D. Chief Executive Officer. DATE: November 13, 2024. By/s/ Michael Kasetaa^a Michael Kasetaa^a Chief Operating Officer and Chief Financial Officer. *91 CERTAIN IDENTIFIED INFORMATION HAS BEEN EXCLUDED FROM THIS EXHIBIT BECAUSE IT IS NOT MATERIAL AND WOULD LIKELY CAUSE COMPETITIVE HARM TO THE REGISTRANT IF PUBLICLY DISCLOSED. *** INDICATES THAT INFORMATION HAS BEEN REDACTED. * Exhibit 10.4*^a * * * * * DEVICE LICENSE AGREEMENT dated as of October 2, 2024 by and between Pharmosa Biopharm Inc. and Liquida Technologies, Inc. * * * * * ARTICLE 1 DEFINITIONS. * * * * * 1 ARTICLE 2 Licenses AND OTHER RIGHTS. * * * * * 2.1 Grant of License to Company. * * * * * 2.2 Grant of Sublicense by Company. * * * * * 2.3 Licensee Technology Transfer. * * * * * 2.4 Manufacturing Technology Transfer. * * * * * 2.5 Procedures for Licensee Technology Transfer. * * * * * 2.6 Second Device Option. * * * * * 2.7 Company Technology Transfer. * * * * * 3.1 DEVELOPMENT, COMMERCIALIZATION and Manufacture OF DEVICES. * * * * * 3.1.1 Development. * * * * * 3.1.2 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* * * * 3.1.357 * * * * * 3.1.358 * * * * * 3.1.359 * * * * * 3.1.360 * * * * * 3.1.361 * * * * * 3.1.362 * * * * * 3.1.363 * * * * * 3.1.364 * * * * * 3.1.365 * * * * * 3.1.366 * * * * * 3.1.367 * * * * * 3.1.368 * * * * * 3.1.369 * * * * * 3.1.370 * * * * * 3.1.371 * * * * * 3.1.372 * * * * * 3.1.373 * * * * * 3.1.374 * * * * * 3.1.375 * * * * * 3.1.376 * * * * * 3.1.377 * * * * * 3.1.378 * * * * * 3.1.379 * * * * * 3.1.380 * * * * * 3.1.381 * * * * * 3.1.382 * * * * * 3.1.383 * * * * * 3.1.384 * * * * * 3.1.385 * * * * * 3.1.386 * * * * * 3.1.387 * * * * * 3.1.388 * * * * * 3.1.389 * * * * * 3.1.390 * * * * * 3.1.391 * * * * * 3.1.392 * * * * * 3.1.393 * * * * * 3.1.394 * * * * * 3.1.395 * * * * * 3.1.396 * * * * * 3.1.397 * * * * * 3.1.398 * * * * * 3.1.399 * * * * * 3.1.400 * * * * * 3.1.401 * * * * * 3.1.402 * * * * * 3.1.403 * * * * * 3.1.404 * * * * * 3.1.405 * * * * * 3.1.406 * * * * * 3.1.407 * * * * * 3.1.408 * * * * * 3.1.409 * * * * * 3.1.410 * * * * * 3.1.411 * * * * * 3.1.412 * * * * * 3.1.413 * * * * * 3.1.414 * * * * * 3.1.415 * * * * * 3.1.416 * * * * * 3.1.417 * * * * * 3.1.418 * * * * * 3.1.419 * * * * * 3.1.420 * * * * * 3.1.421 * * * * * 3.1.422 * * * * * 3.1.423 * * * * * 3.1.424 * * * * * 3.1.425 * * * * * 3.1.426 * * * * * 3.1.427 * * * * * 3.1.428 * * * * * 3.1.429 * * * * * 3.1.430 * * * * * 3.1.431 * * * * * 3.1.432 * * * * * 3.1.433 * * * * * 3.1.434 * * * * * 3.1.435 * * * * * 3.1.436 * * * * * 3.1.437 * * * * * 3.1.438 * * * * * 3.1.439 * * * * * 3.1.440 * * * * * 3.1.441 * * * * * 3.1.442 * * * * * 3.1.443 * * * * * 3.1.444 * * * * * 3.1.445 * * * * * 3.1.446 * * * * * 3.1.447 * * * * * 3.1.448 * * * * * 3.1.449 * * * * * 3.1.450 * * * * * 3.1.451 * * * * * 3.1.452 * * * * * 3.1.453 * * * * * 3.1.454 * * * * * 3.1.455 * * * * * 3.1.456 * * * * * 3.1.457 * * * * * 3.1.458 * * * * * 3.1.459 * * * * * 3.1.460 * * * * * 3.1.461 * * * * * 3.1.462 * * * * * 3.1.463 * * * * * 3.1.464 * * * * * 3.1.465 * * * * * 3.1.466 * * * * * 3.1.467 * * * * * 3.1.468 * * * * * 3.1.469 * * * * * 3.1.470 * * * * * 3.1.471 * * * * * 3.1.472 * * * * * 3.1.473 * * * * * 3.1.474 * * * * * 3.1.475 * * * * * 3.1.476 * * * * * 3.1.477 * * * * * 3.1.478 * * * * * 3.1.479 * * * * * 3.1.480 * * * * * 3.1.481 * * * * * 3.1.482 * * * * * 3.1.483 * * * * * 3.1.484 * * * * * 3.1.485 * * * * * 3.1.486 * * * * * 3.1.487 * * * * * 3.1.488 * * * * * 3.1.489 * * * * * 3.1.490 * * * * * 3.1.491 * * * * * 3.1.492 * * * * * 3.1.493 * * * * * 3.1.494 * * * * * 3.1.495 * * * * * 3.1.496 * * * * * 3.1.497 * * * * * 3.1.498 * * * * * 3.1.499 * * * * * 3.1.500 * * * * * 3.1.501 * * * * * 3.1.502 * * * * * 3.1.503 * * * * * 3.1.504 * * * * * 3.1.505 * * * * * 3.1.506 * * * * * 3.1.507 * * * * * 3.1.508 * * * * * 3.1.509 * * * * * 3.1.510 * * * * * 3.1.511 * * * * * 3.1.512 * * * * * 3.1.513 * * * * * 3.1.514 * * * * * 3.1.515 * * * * * 3.1.516 * * * * * 3.1.517 * * * * * 3.1.518 * * * * * 3.1.519 * * * * * 3.1.520 * * * * * 3.1.521 * * * * * 3.1.522 * * * * * 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* * * * 3.1.579 * * * * * 3.1.580 * * * * * 3.1.581 * * * * * 3.1.582 * * * * * 3.1.583 * * * * * 3.1.584 * * * * * 3.1.585 * * * * * 3.1.586 * * * * * 3.1.587 * * * * * 3.1.588 * * * * * 3.1.589 * * * * * 3.1.590 * * * * * 3.1.591 * * * * * 3.1.592 * * * * * 3.1.593 * * * * * 3.1.594 * * * * * 3.1.595 * * * * * 3.1.596 * * * * * 3.1.597 * * * * * 3.1.598 * * * * * 3.1.599 * * * * * 3.1.600 * * * * * 3.1.601 * * * * * 3.1.602 * * * * * 3.1.603 * * * * * 3.1.604 * * * * * 3.1.605 * * * * * 3.1.606 * * * * * 3.1.607 * * * * * 3.1.608 * * * * * 3.1.609 * * * * * 3.1.610 * * * * * 3.1.611 * * * * * 3.1.612 * * * * * 3.1.613 * * * * * 3.1.614 * * * * * 3.1.615 * * * * * 3.1.616 * * * * * 3.1.617 * * * * * 3.1.618 * * * * * 3.1.619 * * * * * 3.1.620 * * * * * 3.1.621 * * * * * 3.1.622 * * * * * 3.1.623 * * * * * 3.1.624 * * * * * 3.1.625 * * * * * 3.1.626 * * * * * 3.1.627 * * * * * 3.1.628 * * * * * 3.1.629 * * * * * 3.1.630 * * * * * 3.1.631 * * * * * 3.1.632 * * * * * 3.1.633 * * * * * 3.1.634 * * * * * 3.1.635 * * * * * 3.1.636 * * * * * 3.1.637 * * * * * 3.1.638 * * * * * 3.1.639 * * * * * 3.1.640 * * * * * 3.1.641 * * * * * 3.1.642 * * * * * 3.1.643 * * * * * 3.1.644 * * * * * 3.1.645 * * * * * 3.1.646 * * * * * 3.1.647 * * * * * 3.1.648 * * * * * 3.1.649 * * * * * 3.1.650 * * * * * 3.1.651 * * * * * 3.1.652 * * * * * 3.1.653 * * * * * 3.1.654 * * * * * 3.1.655 * * * * * 3.1.656 * * * * * 3.1.657 * * * * * 3.1.658 * * * * * 3

experiments and tests and results of experimentation and testing (including results of research or development), summaries and information contained in submissions to and information from ethical committees, or Regulatory Authorities, and manufacturing process and development information, results and data, whether or not patentable, all to the extent not claimed or disclosed in a patent or patent application; and (b) compositions of matter, assays, animal models and physical, biological or chemical material, including drug substance samples, intermediates of drug substance samples, drug product samples and intermediates of drug product samples. Â The fact that an item is known to the public shall not be taken to exclude the possibility that a compilation including the item, and/or a development relating to the item, is (and remains) not known to the public. Â à€œKnow-Howâ€ means any rights including copyright, database or design rights protecting such Know-How. Â à€œKnow-Howâ€ excludes Patent Rights.1.27â€œLawâ€ or â€œLawsâ€ means all applicable laws, statutes, rules, regulations, ordinances and other pronouncements having the binding effect of law of any Governmental Body.1.28â€œLicensor Know-Howâ€ means all Know-How that is Controlled by Licensor or any of its Affiliates as of the Effective Date, including what is set forth on Schedule 1.28, or at any time thereafter during the Term that is necessary or useful in the Development, manufacture, use, or Commercialization of Devices or Device Products in the Field. Â 1.29â€œLicensorâ€™s Knowledgeâ€ means, with respect to a matter that is the subject of a given representation or warranty of Licensor, the actual knowledge of the executive officers of Licensor, and the vice presidents and senior directors of Licensorâ€™s research and development department, including the individuals set forth in Schedule 1.29, after making reasonable inquiry into the relevant subject matter. 1.30â€œLicensor Patentsâ€ means all Patent Rights that are Controlled by Licensor or any of its Affiliates as of the Effective Date or at any time thereafter during the Term that are necessary or useful for the research, Development, manufacture, use, or Commercialization of Devices or Device Products in the Field. A Listed on Schedule 1.30 are all Licensor Patents existing as of the Effective Date; provided, that Licensor shall update Schedule 1.30 from time-to-time to include any new Patent Rights that come to be Controlled by Licensor or any of its Affiliates at any time during the Term or on following the Effective Date that are necessary or useful for the Development, manufacture, use, or Commercialization of a Device or Device Product. Â 1.31â€œLicensor Technologyâ€ means the Licensor Patents and the Licensor Know-How. Â 1.32â€œMAAâ€ means a Marketing Authorization Application submitted pursuant to the requirements of the FDA, as more fully defined in 21 U.S. C.F.R.â€ 314.3 et seq, a Biologics License Application submitted pursuant to the requirements of the FDA, as more fully defined in 21 U.S. C.F.R. â€ 601, and any equivalent application submitted in any country in the Territory, including all additions, deletions or supplements thereto, and as any and all such requirements may be amended, or supplanted, at any time.1.33â€œManufacturing Costâ€ means the actual and verifiable costs and expenses paid by Licensor to one (1) or more manufacturers or suppliers (including Device Manufacturers) for the manufacture and supply of Devices or other consumables related thereto, including but not limited to, Licensorâ€™s external, Out-of-Pocket Expenses for materials, production, factory overhead, quality control, quality assurance, bulk and finished packaging, transportation and insurance.1.34â€œOut-of-Pocket Expensesâ€ means expenses actually paid by a Party or its Affiliate to any Third Party.1.35â€œPatent Rightsâ€ means: (a) an issued or granted patent, including any extension, supplemental protection certificate, registration, confirmation, reissue, reexamination or renewal thereof; (b) a pending patent application, including any continuation, divisional, continuation-in-part, substitute or provisional application thereof; and (c) all counterparts or foreign equivalents of any of the foregoing issued by or filed in any country or other jurisdiction.1.36â€œPersonâ€ means any natural person, corporation, firm, business trust, joint venture, association, organization, company, partnership or other business entity, or any government or agency or political subdivision thereof.1.37â€œPN1[***] Deviceâ€ means the [***] nebulizer currently under development by Licensor under the [***], including any improvements, enhancements or modifications thereto or derivatives thereof (including any future generation device thereof). A Schedule 1.37 sets forth a description of the PN1[***] Device as it exists as of the Effective Date. Â The [***] Device will include such further development and changes as are made in accordance with Section 3.1.2.1.38â€œPN2[***] Deviceâ€ means the [***] nebulizer currently under development by Licensor under the [***], including any improvements, enhancements or modifications thereto or derivatives thereof (including any future generation device thereof). Â Schedule 1.38 sets forth a description of the PN2[***] Device as it exists as of the Effective Date. The [***] Device will include such further development and changes as are made in accordance with Section 3.1.2. Â 1.39â€œPrice Approvalsâ€ means, in those countries in the Territory where Regulatory Authorities may approve or determine pricing and/or pricing reimbursement for pharmaceutical or biotechnology products, such pricing and/or pricing reimbursement approval or determination.1.40â€œProductâ€ has the meaning set forth in the Product Agreement.1.41â€œJ1.42â€œRegulatory Approvalâ€ means any and all approvals, licenses, registrations, or authorizations of the relevant Regulatory Authority, including Price Approvals, necessary for the Development, manufacture, use, storage, import, transport or Commercialization of a Device or Device Product in a particular country or jurisdiction. Â For the avoidance of doubt, Regulatory Approval to Commercialize a Device or Device Product shall include Price Approval, if required in a particular country or jurisdiction.1.43â€œRegulatory Authorityâ€ means: (a) in the US, the FDA; or (b) in any other jurisdiction anywhere in the world, any regulatory body with similar regulatory authority over pharmaceutical or biotechnology products.1.44â€œRoyaalty Termâ€ has the meaning set forth in the Product Agreement.1.45â€œSublicenseeâ€ means a Person other than an Affiliate of Company to which Company (or its Affiliate) has, pursuant to Section 2.2, granted sublicense rights under any of the Licensed Rights; provided, that â€œSublicenseeâ€ shall exclude distributors and Subcontractors. Â For clarity, the licensee of a Compulsory License shall not be deemed to be a Sublicensee. 1.46â€œTaxâ€ or â€œTaxesâ€ means any federal, state, local or foreign income, gross receipts, license, payroll, employment, excise, severance, stamp, occupation, premium, windfall profits, environmental, customs duties, capital stock, franchise, profits, withholding, social security, unemployment, disability, real property, personal property, sales, use, transfer, registration, value added, alternative or add-on minimum, estimated, or other tax of any kind whatsoever, including any interest, penalty, or addition thereto, whether disputed or not.1.47â€œTerritoryâ€ means all of the countries, jurisdictions and territories in the world, except China (including Hong Kong and Macao), North Korea, the Republic of Korea, Taiwan, Kingdom of Saudi Arabia, United Arab Emirates, Kuwait, Qatar, Oman, Bahrain, Iraq, Egypt, Lebanon, Jordan, Morocco, Algeria, Iran, Tunisia, Sudan, Yemen, Libya, Syria, Turkey, Malaysia, Indonesia, Thailand, Philippines, Singapore, Brunei, Vietnam, Lao, Cambodia and Myanmar. 1.48â€œThird Partyâ€ means any Person other than Licensor, Company or any of their respective Affiliates.1.49â€œThird Party Actionâ€ means any Action made by a Third Party against either Party that claims that a Device or Device Product, or its use, Development, manufacture or Commercialization infringes or misappropriates such Third Partyâ€™s intellectual property rights.1.50â€œThird Party License Agreementâ€ means any agreement entered into by a Party or its Affiliate with a Third Party, or any amendment or supplement thereto, in each case following the Effective Date, whereby royalties, fees or other payments are to be made by a Party or its Affiliate to such Third Party in connection with the grant of rights under intellectual property rights Controlled by such Third Party, which rights are necessary or useful to Develop, manufacture, have made, import, export, use or Commercialize a Device under the Licensed Rights.1.51 â€œUnited Statesâ€ or â€œUSâ€ means the United States of America, its territories and possessions.1.52â€œUSDâ€ or â€œ\$â€ means the lawful currency of the United States.1.53â€œValid Claimâ€ means (a) a claim of an issued and unexpired patent which has not lapsed or been revoked, abandoned or held unenforceable or invalid by a final decision of a court or governmental or supra-governmental agency of competent jurisdiction, unappealable or unappealed within the time allowed for appeal, and which has not been disclaimed, denied or admitted to be invalid or unenforceable through reissue, reexamination or disclaimer or otherwise, or (b) a pending claim of a patent application which patent application has not been pending for more than five (5) years from the date of filing such application and which claim has not lapsed or been cancelled, withdrawn, abandoned or rejected. 1.54Other Terms. Â The definition of each of the following terms is set forth in the section of this Agreement indicated below: Defined TermSectionâ€œActionâ€6.5.2â€œAgreementâ€Preambleâ€œCompany Indemniteesâ€9.2â€œCompany Patentsâ€6.4.5â€œCompany Tech Transfer Materialsâ€2.7â€œCure Periodâ€10.2.2â€œDevelopment Supportâ€3.1.2(c)â€œDevice Manufacturerâ€2.4â€œDisputesâ€11.1â€œEffective Dateâ€Preambleâ€œFirst Amendmentâ€Recitalsâ€ICCâ€11.3.1â€œLicensed Rightsâ€2.1â€œLicensor Development Activitiesâ€3.1.2(a)â€œLicensor Indemniteesâ€9.1â€œLicensor Relevant Actionâ€6.5.2â€œLicensor Technology Transfer Planâ€2.3â€œLossesâ€9.1â€œNon-Specific Licensor Patentsâ€6.4.2â€œOriginal Product Agreementâ€Recitalsâ€œPartyâ€ and â€œPartiesâ€Preambleâ€œProduct Agreementâ€Recitalsâ€œRegulatory Supportâ€4.3â€œRepresentativesâ€3.1.2(c)â€œRight of First Refusalâ€10.6.2â€œRight of First Refusal Notice Periodâ€10.6.2(b)â€œRulesâ€11.3.1â€œSecond Device Optionâ€2.6â€œSpecific Licensor Patentsâ€6.4.1â€œSubcontractorâ€3.4â€œSupply Agreementâ€3.3.1â€œTermâ€10.1â€œARTICLE 2â€œLicenses AND OTHER RIGHTS2.1Grant of License to Company. Â A Subject to the terms and conditions of this Agreement, Licensor hereby grants to Company and its Affiliates (a) an exclusive (even as to Licensor), royalty-bearing right and license (with the right to sublicense, subject to the provisions of Section 2.2) under the Licensor Technology to Develop, have Developed, manufacture, have manufactured, use and Commercialize Devices and Device Products in the Field in the Territory and (b) a non-exclusive right and license (with the right to sublicense, subject to the provisions of Section 2.2) under the Licensor Technology to Develop, have Developed (but not seek MAA), manufacture, have manufactured, use (but not Commercialize) Devices and Device Products in the Field outside the Territory for the sole purpose of exploiting its right and license under clause (a) (clauses (a) and (b) collectively, the â€œLicensed Rightsâ€). A Notwithstanding the scope of the Licensed Rights as set forth above, Company shall not Develop, have Developed, manufacture, have manufactured, use or Commercialize the PN2[***] Device or PN1[***] Device for any product other than the Existing Product. In any event, except as expressly set forth in Sections 2.6 and 3.3.1, Licensor shall not, and shall not permit or authorize any of its Affiliates or sublicensees or any Third Party to, practice or use any Licensor Technology within the scope of the Licensed Rights that has been exclusively licensed to Company under Section 2.1 (a) above. A 2.2Grant of Sublicense by Company. Â Company shall have the right, in its sole discretion, to grant sublicenses, in whole or in part, through multiple tiers, under the Licensed Rights to Third Parties; provided, however, that (a) the granting by Company of a sublicense shall not relieve Company of any of its obligations hereunder; (b) Licensorâ€™s obligations to such Third Party will be no broader than Licensorâ€™s obligations were to Company under this Agreement prior to the grant of such a sublicense, (c) the rights granted to such Third Party under the Licensor Technology will be consistent with the rights granted to Company under Section 2.1 applicable to the scope of the sublicense granted to such Third Party, (d) Company shall provide a copy of each sublicense (and any sub-sublicense) agreement to Licensor within thirty (30) Business Days after execution of such sublicense (subject to reasonable redactions), (e) the terms of each sublicense (and any sub-sublicense) agreement shall be consistent with all applicable terms of this Agreement, and (f) Company remains primarily responsible for the actions or omissions of its Sublicensees. Â In no event shall Company grant a sublicense in whole of the Licensed Rights (including the entire Field and entire Territory) to any single Third Party and/or its Affiliates without the prior written consent of Licensor, such consent not to be unreasonably withheld, conditioned or delayed. 2.3Licensor Technology Transfer. Â As soon as reasonably practicable after the Effective Date and subject to Section 2.5 and the Licensor technology transfer plan (â€œLicensor Technology Transfer Planâ€) set forth in Schedule 2.3, Licensor will transfer to Company, at Licensorâ€™s cost and expense, all Licensor Know-How pursuant to Section 2.5. Â For the avoidance of doubt, nothing in this Agreement shall be in any way interpreted that Licensor is transferring its ownership or proprietary right to Licensor Technology. 2.4Manufacturing Technology Transfer. Â Upon Companyâ€™s request, Licensor shall transfer to Company all Licensor Know-How for the manufacture of Devices and provide reasonable technical assistance to Company, its Affiliates or any Third Party contract manufacturer for Devices (a â€œDevice Manufacturerâ€) at reasonable, mutually agreed upon charges with respect to Companyâ€™s and its Affiliateâ€™s (and any Device Manufacturerâ€™s) receipt, adoption and establishment of the manufacturing process, including: (a) making available a reasonable number of appropriately trained personnel to provide, on a mutually convenient timetable, technical assistance with respect to such transfer, including technical and design details of all equipment used in the process of manufacturing a Device, (b) using Commercially Reasonable Efforts to promptly assist Company and its Affiliates (and any Device Manufacturer) in obtaining all necessary Regulatory Approvals or modifying existing Regulatory Approvals for the manufacture of Devices by Company, its Affiliate or a Device Manufacturer, (c) allowing Company and its Affiliates (and any Device Manufacturer) to cross reference Licensorâ€™s (and its Affiliateâ€™s) regulatory filings (including, but not limited to, a drug master file) and such other regulatory submissions controlled by Licensor (or its Affiliates) applicable to Devices, as the case may be, (d) supplying analytical test methods and other testing Know-How including method validation required to perform release testing or other testing as may be required by the applicable Regulatory Authority, and (e) upon request by Company, providing Company and its Affiliates (and any Device Manufacturer) with appropriate quantities of reference standards related to Product in order to facilitate its testing.2.5Procedures for Licensor Technology Transfer. Â The technology transfers set forth in Section 2.3 and Section 2.4 shall occur in an orderly fashion and in a manner such that the value, usefulness and confidentiality of the transferred Licensor Know-How are preserved in all material respects in accordance with the Licensor Technology Transfer Plan. Â During the Term, Licensor shall provide to Company full and prompt disclosure, but in no event less frequently than semi-annually, of any Licensor Technology that becomes Controlled by Licensor or any of its Affiliates after the Effective Date and that is necessary or useful to Company to conduct its activities or exercise its rights as contemplated hereunder and shall, promptly following such disclosure, transfer to Company such Licensor Know-How. 2.6Second Device Option. Licensor shall, and hereby does, grant to Company and its Affiliates an exclusive and sole right of first option (the â€œSecond Device Optionâ€), at Companyâ€™s election, to include the PN1[***] Device as a Device under this Agreement. Company may exercise the Second Device Option at any time during the Term upon written notice of such exercise to Licensor. Company shall submit to Licensor payment of an exercise fee in the amount [***] to Licensor within [***] days of receipt of an invoice for such exercise fee. Notwithstanding anything in this Agreement to the contrary, following Companyâ€™s exercise of the Second Device Option: (a) Licensor shall retain the right, and such right shall be sublicensable to Third Parties, to Develop, have Developed, manufacture, have manufactured, use and Commercialize the PN1[***] Device in the Field in the Territory under the Licensed Technology, provided that in no event shall Licensor, its Affiliates or their respective sublicensees have the right to Develop, have Developed, manufacture, have manufactured, use or Commercialize the PN2[***] Device in connection with treprostinil in the Territory, 2.7Company Technology Transfer. During the Term and upon Licensorâ€™s reasonable written request (but no more frequently than twice each Calendar Year), Company shall provide to Licensor copies of all technical information, data, reports and regulatory dossiers generated by or on behalf of Company during the Development and Commercialization of the Device and Device Product that are necessary for Licensor to seek Regulatory Approvals for the Device and Device Product in the Field outside the Territory (the â€œCompany Tech Transfer Materialsâ€) at no cost to Licensor. Licensor shall have the right to incorporate, and sublicense such right to any Third Party to which Licensor licenses the right of development, manufacture or commercialization to such Third Party in any country outside of the Territory, any such Company Tech Transfer Materials into its regulatory filings for Regulatory Approvals for the Device and Device Product in the Field outside of the Territory; provided, however, that, notwithstanding any permitted assignment or transfer pursuant to Section 12.2, in no event shall the Company Tech Transfer Materials (including any rights with respect thereto) be assignable, licensable or otherwise transferable to a Third Party, including any Third Party licensee or successor-in-interest to Licensorâ€™s business to which this Agreement relates, that is a Company Competitor (as such term defined in the Original Product Agreement) in the Territory. ARTICLE 3â€œDEVELOPMENT, COMMERCIALIZATION and Manufacture OF DEVICES3.1Development. 3.1.1General. Â Subject to Section 3.1.2, Company shall have the exclusive right, and sole responsibility and decision-making authority, at Companyâ€™s cost and expense, to Develop Devices and Device Products and to conduct (either itself or through its Affiliates, agents, Subcontractors and/or Sublicensees) all Clinical Trials and non-clinical studies Company believes appropriate to obtain Regulatory Approval for Devices and Device Products in the Field in the Territory. Each Party will notify the other, via the joint steering committee under the Product Agreement, if such notifying Party reasonably believes or expects that any Development of a Device or Device Product conducted by or on behalf of such notifying Party would reasonably be expected to have a material adverse effect on the other Partyâ€™s Development of a Device or Device Product, and the joint steering committee will review and discuss such matter. Â 3.1.2Licensor Development and Support. Â (a)Licensor Development. Â Unless and until Company exercises its Second Device Option, Licensor shall, at its sole cost and expense, be responsible for all pre-clinical and manufacturing development of the PN2[***] Device to meet the minimum design requirements for such Device (separately or for Device Products, as applicable) for Company to exploit the Licensed Rights (collectively, the â€œLicensor Development Activitiesâ€). A The minimum design requirements shall include (x) the specifications set forth in Schedule 1.38, (y) any specifications (including changes thereto) that are reasonably necessary for Company to either obtain or maintain Regulatory Approval of a Device or Device Product in the Territory or avoid infringement or misappropriation of any intellectual property right of a Third Party, and (z) any other specifications established by the JSC pursuant to Section 3.1.2(b) (provided the specification is reasonably necessary for Company to either obtain or maintain Regulatory Approval of a Device or Device Product in the Territory or avoid infringement or misappropriation of any intellectual property right of a Third Party).

activities). A Lessor shall perform the Lessor Development Activities in accordance with a development plan established pursuant to Section 3.1.2(b). A If Company exercises its Second Device Option, (i) the Parties shall cooperate to transition all Lessor Development Activities with respect to the PN1[***] Device to Company; and (ii) Lessor shall have no further obligation to perform Lessor Development Activities (and, for clarity, unless otherwise agreed by the Parties in writing, no obligation to perform any Development activities for Company with respect to the PN2[***] Device). Lessor shall collaborate with Company in connection with Lessor's performance of the Lessor Development Activities in accordance with any reasonable direction provided by Company and shall provide Company with regular progress updates on all Lessor Development Activities. A (b)Minimum Design Requirements and Development Plan. A The JSC (as defined in the Product Agreement) under the Product Agreement shall meet from time to time in accordance with Article 3 of the Product Agreement to determine the minimum design requirements and the development plan for the conduct of the Lessor Development Activities for the applicable Device under Section 3.1.2(a). A Accordingly, the Parties hereby agree that the responsibilities of the JSC under Section 3.3 of the Product Agreement shall include (i) discussing, establishing, reviewing, and, as may be applicable from time to time, amending the minimum design requirements and the development plan for the conduct of the Lessor Development Activities for a Device and (ii) monitoring the progress of the Lessor Development Activities against the development plan. From time to time after the first Regulatory Approval of a MAA (as such term is defined in the Original Product Agreement, as amended) for a Device in the United States, Lessor may identify a nebulizer device owned or controlled by Lessor (or otherwise developed by or on behalf of Lessor) and request confirmation from Company that such nebulizer device will not be deemed a future generation of such Device. A In the event that the Parties are unable to agree as to whether such identified device constitutes a future generation of the PN2[***] Device or PN1[***] Device, then either Party shall have the right to escalate such matter to an independent expert, reasonably acceptable to the other Party, with at least ten (10) years of relevant experience in the field of pharmaceutical nebulizers by providing written notice to the other Party. A If the Parties cannot agree on such independent expert within fifteen (15) Business Days of receipt of such notice, each Party shall select one (1) expert, and the two (2) selected experts shall mutually agree upon the third expert, each expert to have the experience set forth in the preceding sentence. A Within ten (10) Business Days from selection of the expert(s), each Party shall deliver to the expert(s) and the other Party its position as to why the identified device does or does not constitute a future generation of the applicable Device and a memorandum in support thereof. A Within ten (10) Business Days after receipt of the other Party's support memorandum, each Party may submit to the expert(s), with a copy to the other Party, a response to the other Party's support memorandum. A Within thirty (30) days following receipt by the expert(s) of the Parties' reply memoranda, the expert(s) will make a determination as to whether the identified device is or is not a future generation of an applicable Device hereunder. A The decision of the expert(s) shall be final and binding. A (c)Lessor Support. A Lessor shall make its employees, consultants, contractors, advisors and agents (a)Representatives that are knowledgeable regarding the Lessor Technology and a Device (including the properties and functions thereof), available to Company for scientific and technical explanations, advice, on-site support (limited to once a year and one (1) week for each on-site support) and meetings with Regulatory Authorities that may reasonably be required by Company (provided Company shall consider in good faith Lessor's requests regarding when such meetings are scheduled) relating to the Development of such Device (the a)Development Support). A The Development Support shall be provided by Lessor free-of-charge during the Term except for reasonable Out-of-Pocket Expenses.

A 3.1.3Acknowledgement. A The Parties acknowledge that Company intends as of the Effective Date to Develop the PN2[***] Device; however, upon exercise of the Second Device Option, Company shall have the right to transition its exercise of its development rights under Section 3.1.1 from the PN2[***] Device to the PN1[***] Device. A For clarity, following its exercise of the Second Device Option, Company shall have the right, but not the obligation, to Develop the PN2[***] Device at its cost and expense. 3.2Commercialization. A Except for Lessor's rights under Section 2.6(a) and Section 2.6(b) to Commercialize Devices in the Field in the Territory under the Licensed Rights other than in connection with treprostinil, Company shall have the exclusive right, and sole responsibility and decision-making authority, to Commercialize Devices in the Field in the Territory itself or through one (1) or more Affiliates or Sublicensees or other Third Parties selected by Company and shall have the sole decision-making authority and responsibility in all matters relating to the Commercialization of Devices in the Field in the Territory. 3.3Manufacturing. A 3.3.1Lessor Supply. A Until either (a) Company and a Device Manufacturer execute a supply agreement governing the manufacture and supply of Devices to Company or (b) Lessor assigns to Company, and Company assumes, a supply agreement, by and between Lessor and a Device Manufacturer, governing the manufacture and supply of Devices (in either case, a Supply Agreement), Lessor shall supply, under the terms of this Agreement, Company with all of its requirements for such Devices (including, without limitation, the full nebulizer kit including the body, consumables and packaging) at a supply price equal to [***] of Manufacturing Cost for non-commercial supply and at a supply price equal to the Manufacturing Cost for commercial supply under the terms and conditions of this Agreement. A Notwithstanding the foregoing, upon Company's written request, the Parties will negotiate and enter into a separate supply agreement for the manufacture and supply of such Devices by Lessor to Company on terms and conditions consistent with the terms and conditions of this Section 3.3.1 and, to the extent reasonably applicable, the terms and conditions of that certain Non-Commercial Supply Agreement, dated as of December 18, 2023, by and between the Parties. A 3.3.2Device Manufacturer Supply. A Upon execution by Company and a Device Manufacturer, or assignment from Lessor to Company, of a Supply Agreement, as the case may be, and in accordance with the terms thereof, Company shall purchase Devices directly from a Device Manufacturer. A Lessor shall cooperate in the transition of its manufacturing and supply responsibility to Company and Device Manufacturer(s) (including performance of a manufacturing technology transfer in accordance with Section 2.4). A Lessor shall use best efforts to assist Company in effectuating a Supply Agreement on commercially reasonable terms, including purchasing Devices at a fixed amount for the Development, to facilitate the development of a business relationship between Company and the Device Manufacturer, and to inform Company of discussions and communications with a Device Manufacturer until such time as such Supply Agreement has been executed between Company and such Device Manufacturer or such Supply Agreement has been assigned by Lessor and assumed by Company. Company shall use Commercially Reasonable Efforts to sign such Supply Agreement with the Device Manufacturer no later than [***] months prior to the first commercial sale of a Device. 3.4Right to Subcontract of Company. A Company may exercise any of its rights, or perform any of its obligations, under this Agreement (including any of the Licensed Rights) by subcontracting the exercise or performance of any portion of such rights and obligations on Company's behalf to a Third Party that has entered into a subcontract agreement to provide services to Company for the purpose of fulfilling Company's obligations hereunder (a Subcontractor); provided that (a) any subcontract granted or entered into by Company as contemplated by this Section 3.4 of the exercise or performance of any portion of the rights or obligations that Company may have under this Agreement shall not relieve Company from any of its obligations under this Agreement, (b) the terms of each subcontract agreement shall be consistent with all applicable terms of this Agreement and (c) Company shall remain primarily responsible for the actions or omissions of its Subcontractors. In no event shall Company subcontract all of its obligations under this Agreement to a single Third Party and/or its Affiliates, other than in connection with a sublicense as permitted pursuant to Section 2.2, without the prior written consent of Lessor, such consent not to be unreasonably withheld, conditioned or delayed. 3.5Trademarks. As between Lessor and Company, Company shall have the sole authority to select trademarks for the Device and shall own all such trademarks. ARTICLE 4 a)REGULATORY MATTERS4.1Regulatory Filings. A As between Company and Lessor, Company shall make, own and maintain all regulatory filings and Regulatory Approvals for the Device in the Territory and the regulatory filings and Regulatory Approvals for the Clinical Trials for the Device or Device Product conducted outside the Territory at its cost after the Effective Date. A Company shall provide a copy of (a) any substantive written communications, notices, or other materials received from any Regulatory Authorities regarding any of the foregoing regulatory filings for Regulatory Approvals and Regulatory Approvals, (b) any substantive written communications with any Regulatory Authority regarding any of the foregoing regulatory filings for Regulatory Approvals, and (c) any proposed significant written communications with any Regulatory Authority regarding any of the foregoing regulatory filings for Regulatory Approval reasonably in advance of submission and, with respect to clause (c), shall consider all of Lessor's comments thereto in good faith. 4.2Communications with Authorities. A Company (or one of its Affiliates or Sublicensees) shall be responsible, and act as the sole point of contact, for communications with all Regulatory Authorities in the Territory in connection with the Development, Commercialization, and manufacturing of Device. A Following the Effective Date, Lessor shall not initiate, with respect to Device, any meetings or contact with any Regulatory Authorities in the Territory without Company's prior written consent. A To the extent Lessor receives any written or oral communication from any Regulatory Authority in the Territory relating to Device, Lessor shall (a) refer such Regulatory Authority to Company, and (b) as soon as reasonably practicable (but in any event within twenty-four (24) hours), notify Company and provide Company with a copy of any written communication received by Lessor or, if applicable, complete and accurate minutes of such oral communication. A At the request of Company, Lessor shall make available to Company, free of charge, a qualified representative who shall, together with the representatives of Company, participate in and contribute to meetings with the Regulatory Authorities with respect to regulatory matters relating to the Lessor Technology. A 4.3Lessor Support in Regulatory Matters. A Lessor shall make its Representatives that are knowledgeable regarding the Lessor Technology or Device available to Company upon Company's request for regulatory explanations, advice and on-site support, that may reasonably be required by Company relating regulatory matters (including preparation and filing for any Regulatory Approvals) for Devices (the a)Regulatory Support). A The Regulatory Support shall be provided by Lessor free-of-charge during the Term. 4.4Adverse Event Reporting. A The Parties agree to comply with any and all Laws that are applicable as of the Effective Date and thereafter during the Term in connection with Device safety data collection and reporting. A If Lessor has or receives any information regarding any Adverse Event which may be related to the use of Device, then Lessor shall provide Company with all such information in English within such reasonable timelines which enable Company to comply with all Laws and relevant regulations and requirements. A Company shall report to Lessor any Adverse Event culminating in death or permanent disability of a patient or subject who is administered Device. A The information exchanged between the Parties pursuant to this Section 4.4 shall be transmitted by e-mail or overnight courier to the following address: A Transmission to Lessor: Weishu LuPharmosa Biopharm Inc. 11F, No. 508, Section 7, Zhongxiao East Road, Nangang District, Taipei City 115, Taiwan Tel: +886-2-2782-7561#107Fax: +886-2-2782-9013Email: Weishu.lu@pharmosa.com.twA Transmission to Company: a Jennifer WeidmanLiquidia Technologies, Inc. 419 Davis Drive, Suite 100Morrisville, NC 27560USAEmail: 919-704-5916Email: jennifer.weidman@liquidia.com a 4.5Safety Data Exchange Agreement. A Without limitation of Section 4.4, the Parties shall, as soon as practical following the Effective Date, negotiate in good faith and enter into a safety data exchange agreement, which shall set forth standard operating procedures governing the collection, investigation, reporting, and exchange of information concerning adverse drug reactions or other adverse events (including Adverse Events) sufficient to permit each Party to comply with its regulatory and other legal obligations within applicable timeframes. 4.6Recalls. A Company shall have the sole right to determine whether and how to implement a recall or other market withdrawal of any Device in the Territory. A Company shall, to the extent allowed by Law and reasonably practicable, provide written notice to Lessor of any such recall or market withdrawal. A Lessor shall take all actions requested by Company in connection with such recall or other market withdrawal. A ARTICLE 5 a)CONSIDERATIONS5.1Consideration. A The Parties acknowledge and agree that the initial fee payable under the First Amendment and the royalties payable on sales of Devices and Device Products under the Product Agreement constitute fair and adequate consideration for the rights granted to Company in this Agreement. 5.2No Double-Counting. A Notwithstanding anything in this Agreement, the Product Agreement (including the First Amendment) or any other agreement between the Parties or their Affiliates, (a) there will be no double counting of any costs or expenses in the calculation of any amounts due from Company or its Affiliates to Lessor or its Affiliates under this Agreement, the Product Agreement or any other agreement between the Parties or their Affiliates, and (b) in no event shall sales (including the calculation of Net Sales under the Product Agreement) of any Device or Product be double-counted under this Agreement, the Product Agreement or any other agreement between the Parties or their Affiliates, including with respect to the calculation of royalties due under any such agreements or the achievement of any threshold for sales milestones (including Sales Milestones under the Product Agreement) under any such agreements. A ARTICLE 6 a)INTELLECTUAL PROPERTY MATTERS6.1Certification Under Drug Price Competition and Patent Restoration Act. A Each Party shall immediately give written notice to the other Party of any certification of which they become aware filed pursuant to 21 U.S.C. Section 355(b)(2)(A) or 21 U.S.C. Section 355(j)(2)(A) (or any amendments or successor statutes thereto) claiming that any Lessor Patents Covering a Device Product, or the manufacture or use of each of the foregoing, are invalid or unenforceable, or that infringement will not arise from the manufacture, use or sale of a product by a Third Party. 6.2Listing of Patents. A Notwithstanding any Lessor Patent prosecution rights of Lessor under this Agreement, Company shall have the sole right to determine which of the Lessor Patents, if any, shall be listed for inclusion in the Approved Drug Product with Therapeutic Equivalence Evaluations pursuant to 21 U.S.C. Section 355, or any successor Law in the United States, together with any comparable Laws in any other country in the Territory. 6.3Further Assurances. A Lessor shall require all of its employees, and use its Commercially Reasonable Efforts to require its contractors and agents, and any Affiliates and Third Parties working on its behalf under this Agreement (and their respective employees, contractors and agents), to assign to Lessor any Lessor Technology. 6.4Patent Prosecution and Maintenance. 6.4.1Specific Lessor Patents. A With respect to Lessor Patents in the Territory that Cover or are directed to only (a) a Device (including the composition of matter, manufacture, or method of use thereof) and (b) treprostinil (or its class of compounds) (a)Specific Lessor Patents, including the Lessor Patents identified as such in Schedule 1.30 (as may be updated by Company from time to time), Company shall have the first right, and the obligation, to file, prosecute (including initiating or defending any reexamination and reissue proceedings) and maintain, using counsel of Lessor's choosing, such Specific Lessor Patents in Lessor's name in the Territory. A Company shall bear all costs and expenses of filing, prosecuting and maintaining Specific Lessor Patents by promptly forwarding to Lessor copies of all official correspondence (including, but not limited to, applications, office actions, and responses) relating thereto. A Lessor shall have the right, and Company shall provide Lessor a reasonable opportunity, to comment on and advise Company as to the conduct of such filing, prosecution and maintenance of Specific Lessor Patents, provided, however, that Company shall have the final decision-making right for all matters associated with such filing, prosecution and maintenance. A At Company's request, Lessor will provide Company with reasonable free-of-charge assistance in prosecuting Specific Lessor Patents to the extent possible, including providing such data in Lessor's Control that is, in Company's reasonable judgment, needed to support the prosecution of a Specific Lessor Patent. A For clarity, (i) any Lessor Patents that satisfy clause (a) and (b) above with respect to the PN2[***] Device shall be deemed Specific Lessor Patents until such time as Company exercises the Second Device Option at which time such Lessor Patents shall be deemed Non-Specific Lessor Patents, and (ii) any Lessor Patents that satisfy clause (a) and (b) above with respect to PN1[***] Device shall be deemed Non-Specific Lessor Patents until such time as Company exercises the Second Device Option at which time such Lessor Patents shall be deemed Specific Lessor Patents. 6.4.2Non-Specific Lessor Patents. A Subject to Section 6.4.1, with respect to all Lessor Patents in the Territory other than Specific Lessor Patents (a)Non-Specific Lessor Patents, as listed in Schedule 1.30 (as may be updated from time to time by Company), Lessor shall have the first right, and the obligation, to file, prosecute (including initiating or defending any reexamination and reissue proceedings) and maintain, using counsel of Lessor's choosing, such Non-Specific Lessor Patents in Lessor's name. A Lessor shall bear all costs and expenses of filing, prosecuting and maintaining Non-Specific Lessor Patents. A Lessor shall keep Company informed of the status of the filing and prosecution of Non-Specific Lessor Patents by promptly forwarding to Company copies of all official material correspondence (including, but not limited to, applications, office actions, and responses) relating thereto. A Company shall have the right, and Lessor shall provide Company a reasonable opportunity, to comment on and advise Lessor as to the conduct of such filing, prosecution and maintenance of Non-Specific Lessor Patents, and Lessor shall incorporate all reasonable comments of Company, provided, however, that Lessor shall have the final decision-making right for all matters associated with such filing, prosecution and maintenance. Notwithstanding the foregoing of this Section 6.4.2, in the event that Lessor or Company wishes to file any continuation or divisional with respect to any Non-Specific Lessor Patent that claims treprostinil as the explicit and sole active pharmaceutical ingredient, then the prosecution and maintenance of any such continuation or divisional shall be governed by Section 6.4.1. 6.4.3Selection Not to File and Prosecute Lessor Patents. A If either Party elects not to file or to continue to prosecute or maintain a Lessor Patent in the Territory where it is permitted to do so pursuant to Sections 6.4.1 and 6.4.2 above, as applicable, or fails to do so after receipt of notice from the other Party, then it shall notify the other Party in writing at least ninety (90) days before any deadline applicable to the filing, prosecution or maintenance of such Lessor Patent, as the case may be, or any other date by which an action must be taken to establish or preserve such Lessor Patent in such country or possession. A In such case, the other Party shall have the right to pursue the filing or support the continued prosecution or maintenance of such Lessor Patent. A A

6.4.4 Patent Term Extension. Notwithstanding any Licensor Patent prosecution rights of Licensor under this Agreement, Company shall be responsible in Licensor's name, for obtaining patent term extensions or supplemental protection certificates or comparable extensions in any other country in the Territory, wherever available for Specific Licensor Patents in the Territory. A Licensor shall provide Company with all relevant information, documentation and assistance in this respect as may reasonably be requested by Company. Any such assistance, supply of information and consultation shall be provided promptly and in a manner that will ensure that all patent term extensions for Specific Licensor Patents are obtained wherever legally permissible, and to the maximum extent available. In the event that any election with respect to obtaining patent term extensions is to be made, Company shall have the right to make such elections, and Licensor shall abide by all such elections.

6.4.5 Ownership; Company Patents. Each Party shall own all right, title, and interest in and to all Know-How developed by such Party or any of its Affiliates or a Third Party on behalf of such Party (and all Patent Rights claiming or covering such Know-How). The Parties shall jointly own and have an undivided one-half interest in and to all Know-How developed jointly by or on behalf of the Parties (and all Patent Rights claiming or covering such Know-How) with respect to Devices, Device Products or Products, and each Party hereby grants to the other Party a perpetual, irrevocable, fully paid-up, worldwide, fully sublicensable, non-exclusive license under its interest in and to such Know-How and Patent Rights. All determinations of inventorship under this Agreement shall be made in accordance with United States patent law. For avoidance of doubt, Company shall own any Know-How and Patent Rights developed by Company or any of its Affiliates or a Third Party on behalf of Company and shall have the right, but not the obligation, to file, prosecute and maintain any such Patent Rights (collectively, "Company Patents"). A Company shall bear all costs and expenses of filing, prosecuting and maintaining Company Patents and Licensor shall have no right, title or interest in or to Company Patents.

6.5 Enforcement. 6.5.1 Notice. (a) If either Party believes that an infringement, unauthorized use, misappropriation or ownership claim or threatened infringement or other such activity by a Third Party with respect to any Licensor Technology, or if a Third Party claims that any Licensor Patent is invalid or unenforceable, in each case in the Territory, the Party possessing such knowledge or belief shall notify the other Party and provide it with details of such infringement or claim that are known by such Party. (b) In the event that Licensor believes that a Company Patent, if any, is being infringed by a Third Party or if a Third Party claims that any Company Patent is invalid or unenforceable, Licensor shall notify Company and provide it with details of such infringement or claim.

6.5.2 Actions. A Company shall have the exclusive right, at its own cost (subject to the indemnity obligations set forth in Section 9.2), to attempt to resolve any infringement or claim, including by filing an infringement suit, defending against such claim or taking other similar action, with respect to a Licensor Patent in the Territory (each, an "Action") and to compromise or settle any such infringement or claim; provided that the compromise or settlement shall require Licensor's prior written consent if the compromise or settlement will have an adverse impact on Licensor's business outside the Territory or ownership of the Licensor Technology, such consent not to be unreasonably withheld, conditioned or delayed. A Company's request, Licensor shall immediately provide Company with all relevant documentation (as may be requested by Company) evidencing that Company is validly empowered by Licensor to take such an Action. A Licensor shall join Company in such Action upon Company's written request. A Licensor shall provide reasonable assistance to Company, at the Company's cost, including providing access to relevant documents and other evidence and making its employees available. All amounts recovered by Company shall be allocated, first, to the costs and expenses of the Parties incurred to enforce the Licensor Patents and, second, to Company (provided that such remaining amounts after deduction of the costs and expenses of the Action shall be deemed Net Sales for royalty calculation purposes under the Product Agreement). In the event that Company does not bring such Action against the Third Party infringer within ninety (90) days of the notice delivered under Section 6.5.1, Licensor may request in writing that Company bring an Action, and Company shall consider such request in good faith. Notwithstanding the foregoing, in the event that a Third Party institutes a re-examination action or inter partes review proceeding or brings an action where the sole relief sought is declaratory judgment, in each case seeking to have a Licensor Patent declared invalid or unenforceable or if the Action involves a Non Specific Licensor Patent (a "Licensor Relevant Action"), and Company does not elect to defend or initiate such Licensor Relevant Action within thirty (30) days following Licensor's request pursuant to the preceding sentence, Licensor or its licensees shall be free to defend or initiate the Licensor Relevant Action, at its own expense, and retain any award or settlement in its entirety. If necessary, Company shall join or be joined as a party to the Licensor Relevant Action, but shall be under no obligation to participate, except to the extent that such participation is required as a result of being named a party to the Licensor Relevant Action. A Company shall offer reasonable assistance in connection therewith, at no charge to Licensor, except for reimbursement of reasonable Out-of-Pocket Expenses.

6.5.3 Company Patents. A Company shall have the sole right and authority, but not the obligation, to enforce Company Patents against any Third Party infringer; provided, however, that Licensor shall provide reasonable assistance to Company with respect thereto, including providing access to relevant documents and other evidence and making its employees available, subject to Company's reimbursement of any Out-of-Pocket Expenses incurred on an on-going basis in providing such assistance.

6.6 Third Party Actions Claiming Infringement. 6.6.1 Notice. If Company becomes aware of any Third Party Action against Company, Company shall promptly notify Licensor thereof in writing, setting for the facts of such claim in reasonable detail.

6.6.2 Right to Defend. As between the Parties, Company shall have the exclusive right, at its sole expense and with counsel of its sole choice, but not the obligation, to defend a Third Party Action described in Section 6.6.1 and to compromise or settle such Third Party Action; provided, however, that Company shall not enter into the settlement, consent judgment or other voluntary disposition of any such Third Party Action without consent by Licensor if the settlement, consent judgment or voluntary disposition will have an adverse impact on Licensor's business outside of the Territory or Licensor Technology or involve the admission of liability on the part of Licensor. A Licensor shall provide reasonable assistance to Company, at the Company's cost (subject to the indemnity obligations set forth in Section 9.2), including providing access to relevant documents and other evidence and making its employees available.

ARTICLE 7 - CONFIDENTIALITY

7.1 Confidentiality Obligations.

A Each Party agrees that, for the Term and for five (5) years thereafter, such Party shall, and shall ensure that its Representatives hold in confidence all Confidential Information disclosed to it by the other Party pursuant to this Agreement, unless such information: (a) is or becomes generally available to the public other than as a result of disclosure by the recipient; (b) is already known by or in the possession of the recipient at the time of disclosure by the disclosing Party; (c) is independently developed by recipient without use of or reference to the disclosing Party's Confidential Information; or (d) is obtained by recipient from a Third Party that has not breached any obligations of confidentiality. The recipient shall not disclose any of the Confidential Information, except to Representatives of the recipient who need to know the Confidential Information for the purpose of performing the recipient's obligations, or exercising its rights, under this Agreement and who are bound by obligations of non-use and non-disclosure substantially similar to those set forth herein. The recipient shall be responsible for any disclosure or use of the Confidential Information by such Representatives. The recipient shall protect Confidential Information using not less than the same care with which it treats its own confidential information, but at all times shall use at least reasonable care. A Each Party shall: (i) implement and maintain appropriate security measures to prevent unauthorized access to, or disclosure of, the other Party's Confidential Information; (ii) promptly notify the other Party of any unauthorized access or disclosure of such other Party's Confidential Information; and (iii) cooperate with such other Party in the investigation and remediation of any such unauthorized access or disclosure.

7.2 Use. Notwithstanding Section 7.1, a Party may use the Confidential Information of the other Party for the purpose of performing its obligations, or exercising its rights, under this Agreement, including for purposes of: (a) filing or prosecuting patent applications, subject to the terms of Section 6.4; (b) prosecuting or defending litigation; (c) conducting pre-clinical studies or Clinical Trials pursuant to this Agreement or the Product Agreement; (d) seeking or maintaining Regulatory Approval of the Device or Device Product; or (e) complying with Law, including securities Law and the rules of any securities exchange or market on which a Party's securities are listed or traded. In addition to the foregoing, Company may, in furtherance of its rights under this Agreement, disclose Confidential Information of Licensor to any Third Party, provided that such Third Party is bound by obligations of confidentiality at least as stringent as the ones herein. In making any disclosures pursuant to this Section 7.2, the disclosing Party shall, where reasonably practicable, give such advance notice to the other Party of such disclosure requirement as is reasonable under the circumstances and will use its Commercially Reasonable Efforts to cooperate with the other Party in order to secure confidential treatment of such Confidential Information required to be disclosed. In addition, in connection with any permitted filing by either Party of this Agreement with any Governmental Body the filing Party shall endeavor to obtain confidential treatment of economic, trade secret information and such other information as may be requested by the other Party, and shall provide the other Party with the proposed confidential treatment request with reasonable time for such other Party to provide comments, and shall include in such confidential treatment request all reasonable comments of the other Party. For the avoidance of doubt and notwithstanding anything in this Agreement to the contrary, in no event may Licensor use or reference any Confidential Information of Company, including any information reported by Company to Licensor in connection with this Agreement, to engage in any Competitive Action (as defined in the Product Agreement).

7.3 Required Disclosure. The recipient may disclose the Confidential Information to the extent required by Law or court order; provided, however, that the recipient promptly provides to the disclosing party prior written notice of such disclosure and provides reasonable assistance in obtaining an order or other remedy protecting the Confidential Information from public disclosure. If the recipient is required to make a disclosure as described in this Section 7.3, the recipient will furnish only that portion of the Confidential Information that is legally required.

7.4 Publications. A Licensor shall not publish any information relating to a Device without the prior written consent of Company (which consent may be withheld or given in Company's sole discretion), unless such information has already been publicly disclosed either prior to the Effective Date or after the Effective Date through no fault of Licensor or otherwise not in violation of this Agreement. A Company shall have the right to make such publications as it chooses, in its sole discretion, without the approval of Licensor. A Licensor shall submit to Company for Company's written approval (which approval be granted or denied in Company's sole discretion) any publication or presentation (including in any seminars, symposia or otherwise) of information related directly or indirectly to the Device for review and approval at least ninety (90) days prior to submission for the proposed date of publication or presentation.

7.5 Press Releases and Disclosure.

7.5.1 Initial Press Release. The proposed joint public announcement by Licensor and Company of the execution of this Agreement is set forth on Schedule 7.5.1 hereto.

7.5.2 Public Disclosures by Licensor. Except as provided in Section 7.5.4, Licensor may not make any subsequent press release or public announcement regarding the terms of this Agreement or any matter covered by this Agreement, including the Development or Commercialization of Devices or Device Products, without the prior written consent of Company.

7.5.3 Public Disclosures by Company. Except as provided in Section 7.5.4, Company may not make any subsequent press release or public announcement regarding the terms of this Agreement; provided, however, that Company shall have the right to make such press releases as it chooses, in its sole discretion, regarding the status of its Development or Commercialization of Devices or Device Products without the approval of Licensor, provided further, that, to the extent practicable, Company shall use Commercially Reasonable Efforts to notify Licensor in advance of any such press release that would reasonably be expected to trigger any securities filing obligations for Licensor.

7.5.4 Exceptions. Notwithstanding the foregoing, either Party shall have the right, without the approval of the other Party, (a) to make securities filings that such Party determines are required under applicable securities laws and regulations (provided, that to the extent practicable, it provides the text of such planned disclosure to the non-disclosing Party no less than two (2) days prior to disclosure, and has used Commercially Reasonable Efforts to incorporate all reasonable comments of the non-disclosing Party regarding such disclosure); and (b) to make disclosures of information that has been previously published or released in accordance with the terms and conditions of this Agreement. Licensor may disclose the terms and conditions of this Agreement to its licensees of devices and device products to clarify the scope of license to Company of the Devices and Device Products, provided that Licensor shall redact all of the financial terms and conditions set forth in this Agreement prior to such disclosure and further provided that Licensor shall provide the text of such planned disclosure to Company no less than two (2) days prior to disclosure and shall incorporate all reasonable comments of Company regarding such disclosure.

ARTICLE 8 - REPRESENTATIONS, WARRANTIES AND COVENANTS

8.1 Representations and Warranties. A Each Party represents and warrants to the other Party that, as of the Effective Date: (a) such Party is duly organized and validly existing under the Laws of the jurisdiction of its incorporation; (b) such Party has taken all action necessary to authorize the execution and delivery of this Agreement and the performance of its obligations under this Agreement; (c) this Agreement is a legal and valid obligation of such Party, binding upon such Party and enforceable against such Party in accordance with the terms of this Agreement, except as enforcement may be limited by applicable bankruptcy, fraudulent conveyance, insolvency, reorganization, moratorium and other laws relating to or affecting creditors' rights generally and by general equitable principles. The execution, delivery and performance of this Agreement by such Party does not conflict with, breach or create in any Third Party the right to accelerate, terminate or modify any agreement or instrument to which such Party is a party or by which such Party is bound, and does not violate any Law of any Governmental Body having authority over such Party; and (d) such Party has all right, power and authority to enter into this Agreement, to perform its obligations under this Agreement.

8.2 Additional Representations and Warranties of Licensor. A Licensor represents and warrants to Company that, as of the Effective Date: (a) no consent by any Third Party or Governmental Body is required with respect to the execution and delivery of this Agreement by Licensor or the consummation by Licensor of the transactions contemplated hereby; (b) no claims have been asserted or threatened by any Person, nor to Licensor's Knowledge, are there any valid grounds for any claim of any such kind, (i) challenging the validity, effectiveness, or ownership of Licensor Technology, and/or (ii) to the effect that the use, reproduction, modification, manufacturing, distribution, licensing, sublicensing, sale or any other exercise of rights in any of Licensor Technology infringes or will infringe on any intellectual property right of any Person; (c) to Licensor's Knowledge, there is no unauthorized use, infringement or misappropriation of any of Licensor Technology by any employee or former employee of Licensor, or any other Third Party in the Territory; (d) the Licensor Patents are subsisting and all registration, renewal, maintenance and other official fees with respect to the Licensor Patents due on or before the date of this Agreement have been paid in full. A Licensor is the sole assignee and owner of each item listed on Schedule 1.30. To Licensor's Knowledge, the Licensor Patents are not the subject of any litigation procedure, discovery process, interference, reissue, reexamination, opposition, appeal proceedings or any other legal dispute; (e) the Licensor Patents (i) constitute all Patent Rights owned or Controlled by Licensor as of the Effective Date that are directly related to, necessary or useful for, or used in, the Development, Regulatory Approval, manufacture, use, marketing, sale, offer for sale, import, export or Commercialization of the PN2[***] Device or PN1[***] Device in the Territory and (ii) listed on Schedule 1.30. A hereto constitute all Patent Rights that are directly related to, necessary or useful for, or used in, the Development, Regulatory Approval, manufacture, use, marketing sale, offer for sale, import, export or Commercialization of the PN2[***] Device or the PN1[***] in the Territory; (f) the Licensor Know-How (i) constitutes all Know-How owned or Controlled by Licensor as of the Effective Date that is directly related to, or are necessary or useful for, the Development, manufacture, use, or Commercialization of the PN2[***] Device or the PN1[***] Device under the Licensed Rights and (ii) to Licensor's Knowledge, constitutes all Know-How that is directly related to, or are necessary or useful for, the Development, manufacture, use or Commercialization of the PN2[***] Device or PN1[***] Device under the Licensed Rights; (g) all of the Licensor Technology is owned by Licensor or its Affiliates and Licensor has not in-licensed, or otherwise obtained any rights, from a Third Party with respect to the PN2[***] Device or PN1[***] Device or the Licensor Technology; (h) Licensor has not licensed to a Third Party the right to develop the PN2[***] Device or PN1[***] Device; (i) no Third Party has filed, pursued or maintained or threatened in writing to file, pursue or maintain any claim, lawsuit, charge, complaint or other action alleging that any Licensor Patent is invalid or unenforceable; (j) to Licensor's Knowledge, Company's and its Affiliates' and Sublicensees' practice and use of the inventions claimed in the Licensor Patents under the Licensed Rights as permitted herein (including the sale, offer for sale, Commercialization or Regulatory Approval of the PN2[***] Device or PN1[***] Device) will not infringe any intellectual property rights of any Third Party; (k) all Representatives of Licensor who have performed any activities on its behalf in connection with Development regarding the PN2[***] Device or PN1[***] Device have assigned to Licensor the whole of their rights in any intellectual property made, discovered or developed by them as a result of such Development, and no Third Party has any rights to any such intellectual property; (l) Licensor has all right, title and interest in and to the Licensor Technology and Licensor Technology is free and clear of any liens, charges, encumbrances or rights of others to possession or use; (m) Licensor has not previously licensed, assigned, transferred, or otherwise conveyed any right, title or interest in and to the Licensor Technology to any Third Party in the Territory, including any rights with respect to the PN2[***] Device or the PN1[***] Device; (n) to Licensor's Knowledge, the Licensor Technology constitutes all of the intellectual property which could reasonably be expected to be necessary or useful for, or used in, the Development, manufacture, Regulatory Approval, import, export, use, marketing, sale, offer for sale or Commercialization of the PN2[***] Device or PN1[***] Device; (o) the PN2[***] Device and PN1[***] Device each fall within the scope of at least one Valid Claim of at least one of the Licensor Patents listed on Schedule 1.30; (p) to Licensor's Knowledge, there is no additional Third Party licenses that have to be taken now or in the future to guarantee freedom-to-operate to Develop, manufacture and Commercialize the PN2[***] Device or PN1[***] Device without any limitation; (q) the Existing Third Party Agreements constitute all agreements that were entered into by Licensor or its Affiliates with Third

Parties for the development or manufacture or supply of the PN2[***] Device or the PN1[***] Device. A Licensee has provided to Company an accurate, true and complete copy of each of the Existing Third Party Agreements, as amended to date, and each of the Existing Third Party Agreements is in full force and effect. A Licensee is not, and to Licensee's Knowledge no other party to any Existing Third Party Agreement is, in breach or default in the performance of its obligations under any of the Existing Third Party Agreements. A Licensee has not received any notice from any Third Party of any breach, default or non-compliance of Licensee under the terms of any of the Existing Third Party Agreements. A There have been no amendments or other modification to any Existing Third Party Agreements, except as have been disclosed to Company in writing; (r) all tangible information and data provided by or on behalf of Licensee to Company on or before the Effective Date in contemplation of this Agreement was and is true, accurate and complete in all material respects, and Licensee has not failed to disclose, or cause to be disclosed, any information or data that would cause the information and data that has been disclosed to be misleading in any material respect; (s) Licensee (and its Affiliates) has not employed or otherwise used in any capacity, and will not employ or otherwise use in any capacity, the services of any Person debarred under any Law, including under Section 21 USC 335a or any foreign equivalent thereof, with respect to the Licensee Technology or a Device; (t) all Development related to the PN2[***] Device and PN1[***] Device prior to the Effective Date has been conducted in accordance with all Laws; and (u) Licensee Covenants. A Licensee covenants to Company that: (a) Licensee shall fulfill all of its obligations, including but not limited to its payment obligations, under each Existing Third Party Agreement; (b) Licensee shall fulfill all of its obligations, including but not limited to its payment obligations, under each Supply Agreement that related to periods prior to the effectuation or assignment to Company of any such agreement; (c) Licensee shall fulfill all of its obligations, including but not limited to its payment obligations, under any Third Party License Agreement; (d) Licensee shall not amend or waive, or take any action or omit to take any action that would alter, any of Licensee's rights under any Existing Third Party Agreement, Third Party License Agreement or Supply Agreement in any manner that adversely affects, or would reasonably be expected to adversely affect, Company's rights and benefits under this Agreement. A Licensee shall promptly notify Company of any default under, termination or amendment of, any Third Party License Agreement or Supply Agreement; and (e) with respect to each Supply Agreement that is to be assigned to Company hereunder, until such time as such Supply Agreement has been assigned to, and assumed by, Company, (i) Licensee shall not amend or terminate such Supply Agreement, or waive, or take any action or omit to take any action that would alter, any of Licensee's rights under any Supply Agreement, and (ii) Licensee shall promptly notify Company of any default under, or termination or amendment of, any Supply Agreement. A In the case of any default by Licensee under a Supply Agreement, Licensee shall provide Company a reasonable opportunity to cure such default. ARTICLE 9. **INDEMNIFICATION AND INSURANCE.** 9.1 Indemnification by Company. A Company shall indemnify, defend and hold Licensee and its Affiliates and each of their respective employees, officers, directors and agents (the "Licensee Indemnitees") harmless from and against any and all liability, damage, loss, cost or expense (including reasonable attorneys' fees) (collectively, the "Losses") to the extent arising out of Third Party claims or suits to the extent arising out of: (a) the Development, sale, offer for sale, import, export and other Commercialization of a Device by or on behalf of Company, its Affiliates or Sublicensees after the Effective Date; (b) Company's gross negligence or willful misconduct; (c) Company's breach of its obligations under this Agreement; or (d) breach by Company of its representations or warranties set forth in Article 8; except, in each case (a)-(d), to the extent such Losses arise out of (i) any activities set forth in Sections 9.2(a)-(d) for which Licensee is obligated to indemnify any Company Indemnitee under Section 9.2 or (ii) any liability for which Licensee is responsible under the Supply Agreement or any other agreement between Licensee and Company. 9.2 Indemnification by Licensee. A Licensee shall indemnify, defend and hold Company and its Affiliates and each of their respective agents, employees, officers and directors (the "Company Indemnitees") harmless from and against any and all Losses to the extent arising out of Third Party claims or suits to the extent arising out of: (a) Licensee's Development, manufacture, use or Commercialization of the Licensee Technology and Devices prior to the Effective Date; (b) Licensee's gross negligence or willful misconduct; (c) Licensee's breach of its obligations under this Agreement; or (d) breach by Licensee of its representations, warranties or covenants set forth in Article 8; except, in each case (a)-(d), to the extent such Losses arise out of any activities set forth in Sections 9.1(a)-(d) for which Company is obligated to indemnify any Licensee Indemnitee under Section 9.1. 9.3 No Consequential Damages. A EXCEPT WITH RESPECT TO EACH PARTY'S INDEMNIFICATION OBLIGATIONS UNDER SECTION 9.1 OR SECTION 9.2, AS APPLICABLE, IN NO EVENT SHALL EITHER PARTY OR ANY OF ITS AFFILIATES BE LIABLE TO THE OTHER PARTY OR ANY OF ITS AFFILIATES FOR SPECIAL, INDIRECT, INCIDENTAL, CONSEQUENTIAL OR PUNITIVE DAMAGES, INCLUDING LOSS OF PROFITS, WHETHER IN CONTRACT, WARRANTY, TORT, NEGLIGENCE, STRICT LIABILITY OR OTHERWISE ARISING OUT OF OR RELATING TO THIS AGREEMENT, THE TRANSACTIONS CONTEMPLATED HEREIN OR ANY BREACH HEREOF. A NOTWITHSTANDING THE FOREGOING, NOTHING IN THIS AGREEMENT SHALL LIMIT EITHER PARTY FROM SEEKING OR OBTAINING ANY REMEDY AVAILABLE UNDER LAW FOR ANY BREACH OF BY THE OTHER PARTY OF ITS CONFIDENTIALITY OBLIGATIONS UNDER ARTICLE 7. 9.4 Notification of Claims; Conditions to Indemnification Obligations. A As a condition to a Party's right to receive indemnification under this Article 9, it shall: (a) promptly notify the other Party as soon as it becomes aware of a claim or suit for which indemnification may be sought pursuant hereto; (b) cooperate, and cause the individual indemnitees to cooperate, with the indemnifying Party in the defense, settlement or compromise of such claim or suit; and (c) permit indemnifying Party to control the defense, settlement or compromise of such claim or suit, including the right to select defense counsel. A In no event, however, may the indemnifying Party compromise or settle any claim or suit in a manner which admits fault or negligence on the part of the indemnified Party or any indemnitee without the prior written consent of the indemnified Party. A Each Party shall reasonably cooperate with the other Party and its counsel in the course of the defense of any such suit, claim or demand, such cooperation to include without limitation using Commercially Reasonable Efforts to provide or make available documents, information and witnesses. A The indemnifying Party shall have no liability under this Article 9 with respect to claims or suits settled or compromised without its prior written consent. 9.5 Insurance. A During the Term, each Party shall obtain and maintain, at its sole cost and expense, insurance (including any self-insured arrangements) in types and amounts, that are reasonable and customary in the United States and Taiwan, as applicable, pharmaceutical and biotechnology industry for companies engaged in comparable activities. A It is understood and agreed that this insurance shall not be construed to limit either Party's liability with respect to its indemnification obligations hereunder. A Each Party will, except to the extent self-insured, provide to the other Party upon request a certificate evidencing the insurance such Party is required to obtain and keep in force under this Section 9.5. ARTICLE 10. **TERM AND TERMINATION.** 10.1 Term and Expiration. A The term of this Agreement (the "Term") shall commence on the Effective Date and, unless earlier terminated as provided in this Article 10, shall continue in full force and effect, on a country-by-country basis until expiration or termination of the Product Agreement for the Existing Product in such country, at which time this Agreement shall expire in its entirety in such country and the terms of Section 10.5.2(a) shall apply. 10.2 Termination upon Material Breach. 10.2.1 Material Breach. A If a Party breaches any of its material obligations under this Agreement, the Party not in default may give to the breaching Party a written notice specifying the nature of the default, requiring it to cure such breach, and stating its intention to terminate this Agreement if such breach is not cured within [***] days. A If such breach is not cured within [***] days after the receipt of such notice, the Party not in default shall be entitled to terminate this Agreement immediately by written notice to the other Party. A For clarity, such material obligations may apply to the performance of either: (a) this Agreement in its entirety, in which case this provision shall apply to the entire Agreement; (b) a specific Device or Device(s), in which case this provision shall apply only to such affected Device or Device(s); or (c) a specific country or countries within the Territory, in which case this provision shall apply only to such affected country or countries. A For the avoidance of doubt, in no event shall a breach of any material obligation under this Agreement be deemed a breach of any material obligation under the Product Agreement or be deemed to give rise to a right of termination under the Product Agreement. A Termination of this Agreement shall not be deemed to result in termination of the Product Agreement. 10.2.2 Licensee Cure Period. A If Licensee is the defaulting party and a material breach by Licensee is not cured within [***] days of receipt following a notice from Company under Section 10.2.1 (the "Cure Period"), Company may elect not to terminate this Agreement and, instead, during the period commencing at the end of the Cure Period and continuing until the end of the last Royalty Term in all countries, reduce the Development Milestone payments under Section 6.2 of the Product Agreement, the Sales Milestone payments under Section 6.3 of the Product Agreement and the then-applicable royalty rates under Section 6.4.1 of the Product Agreement by [***]; provided, that such reduction shall not be Company's sole remedy with respect to the breach by Licensee. 10.2.3 Material Breach Dispute. A Any Dispute regarding an alleged material breach of this Agreement shall be resolved in accordance with Article 11. A In such event, termination will be tolled and the termination will become effective only if such material breach remains uncured for the applicable cure period after the final resolution of the Dispute through such dispute resolution procedures. 10.3 Bankruptcy Event Termination. A This Agreement may be terminated by written notice by a Party at any time during the Term in the event of a Bankruptcy Event of the other Party. 10.4 Mutual Termination. A The Parties may terminate this Agreement in its entirety or on a country-by-country or Device-by-Device basis upon mutual written agreement. 10.5 Effects of Termination. 10.5.1 Survival. (a) Notwithstanding the expiration or termination of this Agreement, the following provisions shall survive: A Articles 1, 7 (solely with respect to the time period set forth in Section 7.1) and 11; and Sections 3.5 (with respect to trademark ownership), 4.1 (with respect to ownership of regulatory filings and Regulatory Approvals), 4.6, 5.2, 6.4.5, 9.1-9.4, 10.5-10.7, 12.1, 12.2.1-12.2.4, 12.2.5 (for so long as Company has a continuing license hereunder), 12.3, and 12.4-12.17. (b) Expiration or termination of this Agreement shall not relieve the Parties of any liability that accrued hereunder prior to the effective date of such termination. A In addition, termination of this Agreement shall not preclude either Party from pursuing all rights and remedies it may have hereunder or at Law or in equity with respect to any breach of this Agreement nor prejudice either Party's right to obtain performance of any obligation. 10.5.2 Licenses. (a) As of the effective date of expiration of the Term with respect to a given country, the Licensed Rights shall convert to a fully paid, royalty free, irrevocable, perpetual, exclusive, and sublicensable license under the Licensee Technology to Develop, manufacture, have manufactured, use and Commercialize Devices and Device Products in the Field in such country. (b) Upon termination of this Agreement by Licensee pursuant to Section 10.2.1 or 10.3, the following terms and conditions shall apply with respect to such Device(s) or Device Product(s) and country(ies) as are the subject of such termination: (i) all licenses granted to Company under Section 2.1 shall terminate; (ii) Company shall, upon written request by Licensee and within three (3) months therefrom, and subject to Licensee assuming legal responsibility for any Clinical Trials of a Device or Device Product then ongoing, transfer to Licensee or its Third Party designee at no cost to Licensee (except in any such case where Licensee is seeking a claim for damages from Company with respect to any such breach or termination of this Agreement) ownership and control of all regulatory filings, Regulatory Approvals and Device or Device Product data prepared or obtained by or on behalf of Company prior to the date of such termination, to the extent solely related to the Device or Device Product and country(ies) and transferable, and Company shall take any actions reasonably necessary to effect such transfer, provided Company shall have the right to retain one copy of such transferred regulatory filings, Regulatory Approvals and Device or Device Product data for record-keeping purposes; (iii) Company shall, upon written request of Licensee, return to Licensee or, at Licensee's option, destroy, at Licensee's cost and expense, all relevant records and materials in its possession or control containing or comprising the Licensee Know-How, or such other Confidential Information of Licensee, to the extent solely related to such Device(s) or Device Product(s) and country(ies); provided, however, that Company shall have the right to retain one copy of such Licensee Know-How and such other Confidential Information of Licensee for archival purpose; (iv) Company shall, at Licensee's election within thirty (30) days following termination, sell such materials (in whole or in part) to Licensee at a price equal to Company's costs of goods. A Any clinical supplies of such Device(s), Device Product(s) or other materials purchased by Licensee from Company shall be purchased on an à la carte basis with no representations or warranties. A In the event that Licensee does not make an election within such thirty (30) day period or elects not to purchase such materials, Company shall have the right to (A) destroy or retain any and all chemical, biological or physical materials relating to or comprising such Device(s) or Device Product(s), including clinical supplies of such Device(s) or Device Product(s), that are Controlled by Company to the extent solely related to such country(ies) or (B) sell such materials to a Third Party; (v) To the extent not prohibited by Law, Company shall wind down any ongoing Clinical Trials to the extent solely related to such Device(s) or Device Product(s) and country(ies); (vi) Company and its Affiliates and Sublicensees shall be entitled, during the [***] month period following such termination, so long as Company pays to Licensee the royalties applicable to said subsequent sales in accordance with the terms and conditions set forth in this Agreement. A Any commercial inventory remaining following [***] month period shall be offered for sale to Licensee at a price equal to Company's costs of goods; and (vii) Upon any termination of this Agreement, each of Company's Sublicensees shall continue to have the rights and license set forth in its sublicense agreements, which agreements shall be automatically assigned to Licensee, to the extent solely related to such Device(s) or Device Product(s) and country(ies); provided, however, that such Sublicensee is not then in breach of any of its material obligations under its sublicense agreement. 10.6 Additional Effects of Termination for a Licensee Bankruptcy Event. 10.6.1 Continuing Rights. A The Parties agree that Company, as licensee of such rights under this Agreement, shall retain and may fully exercise all of its rights and elections under the U.S. Bankruptcy Code. A The Parties further agree that, in the event of a Licensee Bankruptcy Event, Company shall be entitled to a complete duplicate of (or complete access to, as appropriate) any such intellectual property and all embodiments of such intellectual property, which, if not already in Company's possession, shall be promptly delivered to it (a) following any such commencement of a bankruptcy proceeding upon Company's written request therefor, unless Licensee elects to continue to perform all of its obligations under this Agreement or (b) if not delivered under clause (a), following the rejection of this Agreement by Licensee upon written request therefor by Company. 10.6.2 Right of First Refusal. A In addition to the foregoing, in the event of a Licensee Bankruptcy Event, Company shall, to the extent allowed by Law (including to the extent enforceable under the Laws of Taiwan), have a right of first refusal to purchase all of Licensee's interest in the Device or Device Product and the Licensee Technology (the "Right of First Refusal"). A The Right of First Refusal shall operate as follows: (a) Licensee (or other authorized representative of Licensee, including a bankruptcy trustee) shall promptly send to Company a reasonably detailed written notification of any Licensee Bankruptcy Event. (b) Licensee (or other authorized representative of Licensee, including a bankruptcy trustee) shall promptly send to Company a written notification of any Third Party offer made on Device, Device Product or Licensee Technology. A For a period of up to [***] days after Company receives such notice (such period, the "Right of First Refusal Notice Period"), it shall notify Licensee of its intention to exercise its Rights of First Refusal. A In the event Company exercises its Right of First Refusal, the terms of the Third Party offer shall become binding upon Company and Licensee. A For the avoidance of doubt, Licensee shall not enter into any agreement with a Third Party relating to Licensee's interest in the Devices, Device Products or Licensee Technology during the Right of First Refusal Notice Period. 10.7 Other Remedies. A Termination of this Agreement for any reason shall not release either Party from any liability or obligation that already has accrued prior to such termination. A Termination of this Agreement for any reason shall not constitute a waiver or release of, or otherwise be deemed to prejudice or adversely affect or limit, any rights or remedies that otherwise may be available at Law or in equity. ARTICLE 11. **DISPUTE RESOLUTION.** 11.1 General. A The Parties recognize that disputes (à la carte Disputes) as to certain matters may from time to time arise during the Term which relate to either Party's rights and/or obligations hereunder. A It is the objective of the Parties to establish under this Article 11 procedures to facilitate the resolution of Disputes arising under this Agreement in an expedient manner by mutual cooperation and without resort to litigation. A 11.2 Escalation to Executive Officers. A Either Party may, by written notice to the other Party, request that a Dispute that remains unresolved by the Parties for a period of thirty (30) days be submitted to the Executive Officers for resolution. A If the Executive Officers cannot resolve such Dispute within thirty (30) days after referral of such Dispute to them, then, at any time after such thirty (30) day period, either Party may refer such Dispute to arbitration by submitting a written notice of such request to the other Party. A 11.3 Arbitration. A The Parties hereby agree that, except as otherwise expressly set forth herein, in the event the Parties are unable to resolve any Dispute after referring such Dispute to the Executive Officers, the Dispute shall be settled by binding arbitration administered by the International Chamber of Commerce (à la carte ICC) in accordance with its Rules of Arbitration (the "Arbitration Rules"). A Either Party may refer any Dispute to arbitration by submitting a written notice of such request to the other Party. 11.3.2 Arbitrators. A Any arbitration shall be presided over by three (3) arbitrators. A Each Party shall select one (1) arbitrator, and such selected arbitrators shall mutually agree upon the third arbitrator who shall act as the chairman of the arbitration panel. A If either Party fails or both Parties fail to choose an arbitrator or arbitrators within thirty (30) days after receiving notice of commencement of arbitration or if the two (2) arbitrators fail to choose a third arbitrator within thirty (30) days after their appointment, then either or both Parties shall immediately request that the ICC select the remaining number of arbitrators to be selected. A The arbitrators shall be neutral and independent of the Parties and their respective Affiliates, and may not be current or former directors, officers or employees of the Parties or their respective Affiliates. A No Party may have any ex parte discussion with any potential arbitrator, except for confirming if such arbitrator is willing and able to serve on the arbitration panel. A All arbitrators shall have ten (10) or more years of experience in the pharmaceutical and biotechnology

process, summarize and report financial information; andâ€¢**(b)**Any fraud, whether or not material, that involves management or other employees who have a significant role in the registrantâ€™s internal control over financial reporting.â€¢â€¢â€¢**Date:** November 13, 2024By:/s/ Roger A. Jeffs, Ph.D.â€¢**Name:**Roger A. Jeffs, Ph.D.â€¢**Title:**Chief Executive Officerâ€¢â€¢**(Principal Executive Officer)**â€¢**Exhibit 31.2â€¢CERTIFICATION OF THE PRINCIPAL FINANCIAL OFFICERPURSUANT TO SECTION 302 OF THE SARBANES-OXLEY ACT OF 2002â€¢**i, Michael Kasetta, certify that:â€¢**1.**I have reviewed this Quarterly Report on Form 10-Q of Liquidia Corporation;â€¢**2.**Based on my knowledge, this report does not contain any untrue statement of a material fact or omit to state a material fact necessary to make the statements made, in light of the circumstances under which such statements were made, not misleading with respect to the period covered by this report;â€¢**3.**Based on my knowledge, the financial statements, and other financial information included in this report, fairly present in all material respects the financial condition, results of operations and cash flows of the registrant as of, and for, the periods presented in this report;â€¢**4.**The registrantâ€™s other certifying officer and I are responsible for establishing and maintaining disclosure controls and procedures (as defined in Exchange Act Rules 13a-15(e) and 15d-15(e)) and internal control over financial reporting (as defined in Exchange Act Rules 13a-15(f) and 15d-15(f)) for the registrant and have:â€¢**(a)**Designed such disclosure controls and procedures, or caused such disclosure controls and procedures to be designed under our supervision, to ensure that material information relating to the registrant, including its consolidated subsidiaries, is made known to us by others within those entities, particularly during the period in which this report is being prepared;â€¢**(b)**Designed such internal control over financial reporting, or caused such internal control over financial reporting to be designed under our supervision, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with generally accepted accounting principles;â€¢**(c)**Evaluated the effectiveness of the registrantâ€™s disclosure controls and procedures and presented in this report our conclusions about the effectiveness of the disclosure controls and procedures, as of the end of the period covered by this report based on such evaluation; andâ€¢**(d)**Disclosed in this report any change in the registrantâ€™s internal control over financial reporting that occurred during the registrantâ€™s most recent fiscal quarter (the registrantâ€™s fourth fiscal quarter in the case of an annual report) that has materially affected, or is reasonably likely to materially affect, the registrantâ€™s internal control over financial reporting; andâ€¢**5.**The registrantâ€™s other certifying officer and I have disclosed, based on our most recent evaluation of internal control over financial reporting, to the registrantâ€™s auditors and the audit committee of the registrantâ€™s board of directors (or persons performing the equivalent functions);â€¢**(a)**All significant deficiencies and material weaknesses in the design or operation of internal control over financial reporting which are reasonably likely to adversely affect the registrantâ€™s ability to record, process, summarize and report financial information; andâ€¢**(b)**Any fraud, whether or not material, that involves management or other employees who have a significant role in the registrantâ€™s internal control over financial reporting.â€¢â€¢â€¢â€¢**Date:** November 13, 2024By:/s/ Michael Kasettaâ€¢**Name:**Michael Kasettaâ€¢**Title:**Chief Financial Officer and Chief Operating Officerâ€¢â€¢**(Principal Financial Officer)**â€¢**Exhibit 32.1â€¢CERTIFICATION OF THE PRINCIPAL EXECUTIVE OFFICERPURSUANT TO SECTION 906 OF THE SARBANES-OXLEY ACT OF 2002â€¢**In connection with the Quarterly Report of Liquidia Corporation, a Delaware corporation (the â€œCompanyâ€), on Form 10-Q for the nine months ended September 30, 2024, as filed with the Securities and Exchange Commission on the date hereof (the â€œReportâ€), I, Roger A. Jeffs, Ph.D., Chief Executive Officer of the Company, hereby certify, pursuant to 18 U.S.C. â 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002, that to my knowledge:â€¢**1.**The Report fully complies with the requirements of Section 13(a) or 15(d) of the Securities Exchange Act of 1934, as amended; andâ€¢**2.**The information contained in the Report fairly presents, in all material respects, the financial condition and results of operations of the Company.â€¢â€¢â€¢â€¢**Date:** November 13, 2024By:/s/ Roger A. Jeffs, Ph.D.â€¢**Name:**Roger A. Jeffs, Ph.D.â€¢**Title:**Chief Executive Officerâ€¢â€¢**(Principal Executive Officer)**â€¢**Exhibit 32.2â€¢CERTIFICATION OF THE PRINCIPAL FINANCIAL OFFICERPURSUANT TO SECTION 906 OF THE SARBANES-OXLEY ACT OF 2002â€¢**In connection with the Quarterly Report of Liquidia Corporation, a Delaware corporation (the â€œCompanyâ€), on Form 10-Q for the nine months ended September 30, 2024, as filed with the Securities and Exchange Commission on the date hereof (the â€œReportâ€), I, Michael Kasetta, Chief Financial Officer of the Company, hereby certify, pursuant to 18 U.S.C. â 1350, as adopted pursuant to Section 906 of the Sarbanes-Oxley Act of 2002, that to my knowledge:â€¢**1.**The Report fully complies with the requirements of Section 13(a) or 15(d) of the Securities Exchange Act of 1934, as amended; andâ€¢**2.**The information contained in the Report fairly presents, in all material respects, the financial condition and results of operations of the Company.â€¢â€¢â€¢â€¢**Date:** November 13, 2024By:/s/ Michael Kasettaâ€¢**Name:**Michael Kasettaâ€¢**Title:**Chief Financial Officer and Chief Operating Officerâ€¢â€¢**(Principal Financial Officer)**â€¢