HORIZON PHARMA, INC. Form DEFA14A June 23, 2014

## **UNITED STATES**

## **SECURITIES AND EXCHANGE COMMISSION**

Washington, D.C. 20549

## **SCHEDULE 14A**

#### (RULE 14a-101)

#### **SCHEDULE 14A INFORMATION**

Proxy Statement Pursuant to Section 14(a) of the

Securities Exchange Act of 1934

Filed by the Registrant x

Filed by a Party other than the Registrant "

Check the appropriate box:

- " Preliminary Proxy Statement
- " Confidential, for Use of the Commission Only (as permitted by Rule 14a-6(e)(2))
- " Definitive Proxy Statement
- " Definitive Additional Materials
- x Soliciting Material Pursuant to 240.14a-12

# Horizon Pharma, Inc.

(Name of Registrant as Specified In Its Charter)

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#### (Name of Person(s) Filing Proxy Statement, if other than the Registrant)

Payment of Filing Fee (Check the appropriate box):

- x No fee required.
- " Fee computed on table below per Exchange Act Rules 14a-6(i)(1) and 0-11.
  - (1) Title of each class of securities to which transaction applies:
  - (2) Aggregate number of securities to which transaction applies:
  - (3) Per unit price or other underlying value of transaction computed pursuant to Exchange Act Rule 0-11 (set forth the amount on which the filing fee is calculated and state how it was determined):
  - (4) Proposed maximum aggregate value of transaction:
  - (5) Total fee paid:
- " Fee paid previously with preliminary materials.
- " Check box if any part of the fee is offset as provided by Exchange Act Rule 0-11(a)(2) and identify the filing for which the offsetting fee was paid previously. Identify the previous filing by registration statement number, or the Form or Schedule and the date of its filing.
  - (1) Amount Previously Paid:
  - (2) Form, Schedule or Registration Statement No.:
  - (3) Filing Party:

(4) Date Filed:

Filed under Rule 14a-12

of the Securities Exchange Act of 1934

Filing by: Horizon Pharma, Inc.

Subject Company: Horizon Pharma, Inc.

SEC File No. of Horizon Pharma, Inc.: 001-35238

This Schedule 14A filing consists of a presentation that will be used by Horizon Pharma, Inc. ( Horizon ) in investor meetings and conferences beginning on June 23, 2014. Certain information contained in the presentation relating to Vidara Therapeutics International Ltd. ( Vidara ) and ACTIMMU®E been provided by Vidara.

#### **Forward Looking Statements**

The presentation contains forward-looking statements, including, but not limited to, statements related to the anticipated consummation of a business combination transaction between Horizon and Vidara and the timing and benefits thereof, Horizon s and the combined company s strategy, plans, objectives, expectations (financial or otherwise) and intentions, future financial results and growth potential, anticipated product portfolio, development programs and management structure, and other statements that are not historical facts. These forward-looking statements are based on Horizon s current expectations and inherently involve significant risks and uncertainties. Actual results and the timing of events could differ materially from those anticipated in such forward looking statements as a result of these risks and uncertainties, which include, without limitation, risks related to Horizon s ability to complete the transaction with Vidara on the proposed terms and schedule; risks associated with business combination transactions, such as the risk that the businesses will not be integrated successfully, that such integration may be more difficult, time-consuming or costly than expected or that the expected benefits of the transaction will not occur; risks related to future opportunities and plans for Horizon, as well as the combined company, including uncertainty of the expected financial performance and results; disruption from the proposed transaction, making it more difficult to conduct business as usual or maintain relationships with customers, employees or suppliers; the calculations of, and factors that may impact the calculations of, the acquisition price in connection with the proposed merger and the allocation of such acquisition price to the net assets acquired in accordance with applicable accounting rules and methodologies; and the possibility that if the combined company does not achieve the perceived benefits of the proposed transaction as rapidly or to the extent anticipated by financial analysts or investors, the market price of the combined company s shares could decline, as well as other risks related to Horizon s business, including Horizon s dependence on sales of DUEXIS and VIMOVO and its ability to increase sales of its DUEXIS, VIMOVO and RAYOS/LODOTRA products; competition, including potential generic competition; the ability of Horizon to protect its intellectual property and defend its patents; regulatory obligations and oversight; and those risks detailed from time-to-time under the caption Risk Factors and elsewhere in Horizon s SEC filings and reports, including in its Annual Report on Form 10-K for the year ended December 31, 2013. Horizon undertakes no duty or obligation to update any forward-looking statements contained in this presentation as a result of new information, future events or changes in its expectations.

#### Additional Information and Where to Find It

In connection with the proposed transaction with Vidara, Horizon and Vidara will be filing documents with the SEC, including the filing by Horizon of a preliminary and definitive proxy statement/prospectus relating to the proposed transaction and the filing by Vidara of a registration statement on Form S-4 that will include the proxy statement/prospectus relating to the proposed transaction. After the registration statement has been declared effective by the SEC, a definitive proxy statement/prospectus will be mailed to Horizon stockholders in connection with the

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proposed transaction. INVESTORS AND SECURITY HOLDERS ARE URGED TO READ THE REGISTRATION STATEMENT ON FORM S-4 AND THE RELATED PRELIMINARY AND DEFINITIVE PROXY/PROSPECTUS WHEN THEY BECOME AVAILABLE BECAUSE THEY WILL CONTAIN IMPORTANT INFORMATION ABOUT HORIZON, VIDARA AND THE PROPOSED TRANSACTION. Investors and security holders may obtain free copies of these documents (when they are available) and other related documents filed with the SEC at the SEC s web site at www.sec.gov, by directing a request to Horizon s Investor Relations department at Horizon Pharma, Inc., Attention: Investor Relations, 520 Lake Cook Road, Suite 520, Deerfield, IL 60015 or to Horizon s Investor Relations department at 224-383-3000 or by email to investor-relations@horizonpharma.com. Investors and security holders may obtain free copies of the documents filed with the SEC on Horizon s website at www.horizonpharma.com under the heading Investors and then under the heading SEC Filings.

Horizon and its directors and executive officers and Vidara and its directors and executive officers may be deemed participants in the solicitation of proxies from the stockholders of Horizon in connection with the proposed transaction. Information regarding the special interests of these directors and executive officers in the proposed transaction will be included in the proxy statement/prospectus described above. Additional information regarding the directors and executive officers of Horizon is also included in Horizon s Annual Report on Form 10-K for the year ended December 31, 2013, which was filed with the SEC on March 13, 2014. These documents are available free of charge at the SEC s web site at www.sec.gov and from Investor Relations at Horizon as described above.

This communication does not constitute an offer to sell, or the solicitation of an offer to sell, or the solicitation of an offer to subscribe for or buy, any securities nor shall there be any sale, issuance or transfer of securities in any jurisdiction in which such offer, solicitation or sale would be unlawful prior to registration or qualification under the securities laws of any such jurisdiction.

NASDAQ: HZNP June 2014 Filed under Rule 14a-12 of the Securities Exchange Act of 1934 Filing by: Horizon Pharma, Inc. Subject Company: Horizon Pharma, Inc. SEC File No. of Horizon Pharma, Inc.: 001-35238 The following is a slide presentation relating to the proposed transactions

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described therein that was made available beginning on June 23, 2014. Horizon Pharma, Inc.

Forward-Looking Statements

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Additional Information

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Horizon Pharma and its directors and executive officers and Vidara Therapeutics and its directors and executive officers may be deemed participants in the solicitation of proxies from the stockholders of Horizon Pharma in connection with the proposed transaction. Information regarding the special interests of these directors and executive officers in the proposed transaction will be included in the proxy statement/prospectus described above. Additional information regarding the directors and executive officers of Horizon Pharma is also included in Horizon Pharma s Annual Report on Form 10-K for the year ended December 31, 2013, which was filed with the SEC on March 13, 2014. These documents are available free of charge at the SEC s web site at www.sec.gov and from Investor Relations at Horizon Pharma as described above. This communication does not constitute an offer to sell, or the solicitation of an offer to sell, or the solicitation of an offer to subscribe for or buy, any securities nor shall there be any sale, issuance or transfer of securities in any jurisdiction in which such offer, solicitation or sale would be unlawful prior to registration or qualification under the securities laws of any such jurisdiction.

For full prescribing information refer to product websites.

Note Regarding Use of Non-GAAP Financial Measures

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Horizon Pharma provides non-GAAP net income (loss) and net income (loss) per share financial measures that include adjusting GAAP figures. These adjustments to GAAP exclude non-cash items such as stock compensation and depreciation and amortiz cash interest expense, and other non-cash charges. Certain one-time or substantive events may also be included in the non-GA adjustments periodically when their magnitude is significant within the periods incurred. EBITDA, or earnings before interest depreciation and amortization, is also used and provided by Horizon Pharma as a non-GAAP financial measure. Horizon Pharma examples are appreciated as a non-GAAP financial measure.

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that these non-GAAP financial measures, when considered together with the GAAP figures, can enhance an overall understand Horizon Pharma s financial performance. The non-GAAP financial measures are included with the intent of providing investor more complete understanding of operational results and trends. In addition, these non-GAAP financial measures are among the indicators Horizon Pharma s management uses for planning and forecasting purposes and measuring Horizon Pharma s perfor These non-GAAP financial measures should be considered in addition to, and not as a substitute for, or superior to, financial measures calculated in accordance with GAAP. The non-GAAP financial measures used by Horizon Pharma may be calculated different therefore may not be comparable to, non-GAAP financial measures used by other companies.

(1)
On a non-GAAP basis
(2)
Pending the closing of the acquisition of Vidara Therapeutics International Ltd. which is expected this summer
(3)

RAYOS is known as LODOTRA outside the United States Profitable

(1)

, specialty pharma company with accelerating growth
Integrated commercial model with analytics as its foundation
Four
products
targeting
unmet
therapeutic
needs
in
primary
care,
orphan
diseases
(2)
and specialty segments

#### VIMOVO

R (naproxen/esomeprazole)

DUEXIS ® (ibuprofen/famotidine)

ACTIMMUNE ®

(interferon gamma 1b) (2)

#### RAYOS

(prednisone) delayed-release tablets
(3)
Tax efficient corporate platform facilitating an aggressive business development strategy via product/company acquisitions
(2)
Proven leadership team
5
Horizon Pharma Overview

Accelerating Growth in Revenues and EBITDA 6 ~497% Year-over-Year Net Sales Growth 2011 2012 2013

2014E
(1)
1Q 2013
1Q 2014
\$6.9
\$18.8
\$74.0
\$275.0
\$8.7
\$51.9
\$(46.8)
\$(73.3)
\$(33.5)
\$(16.6)
\$11.0
\$(150.0)
\$(100.0)
\$(50.0)
\$-
\$50.0
\$100.0
\$150.0
\$200.0
\$250.0
\$300.0
Net Sales
Adjusted EBITDA
(1)
Midpoint of 2014 guida
1 1 1 0 1

Midpoint of 2014 guidance provided on May 9, 2014 for net sales of \$270 - \$280 million and adjusted EBITDA of \$80 - \$90 assumed period of August through December 2014 and excluded transaction related expenses. By this presentation, Horizon is guidance. \$85.0

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Integrated Commercial Model Leading-Edge, Value-Based Analytics Differentiated Sales Approach Do What is Right for the Patient Optimize Value Prescriptions Made Easy

#### Rep profile B2B

Funnel management

Optimized targeting

Total office and pharmacy call

Uncapped incentives

\$0 co-pay program

Ensure ubiquity

Align WAC and co-pay

Maximize net revenues

Understand the interplay of pricing, managed care control and script volume

Eliminate script fulfillment friction

Specialty pharmacy channel

HZNP guarantees reimbursement

Integrated Commercial Model *(continued)* DUEXIS Unique Prescribers and Adopters Continue to Grow 8 Added 200+ new writers every week for last 20 months (1) Source: IMS Xponent data Number of Unique Writers (1)
 13% increase over last 3 months
 Number of Unique Adopters (5+ TRx)
 (1)
 19% increase over last 3 months
 Unique Prescribers

+13%

Unique Adopters (5+ Rx/week)

+19%

~35% of DUEXIS Prescriptions Through PME (May 2014) Rx Filled Refill Rate (May 2014) (2) Fill Rate (1)
9
Prescriptions-Made-Easy
(PME) Specialty
Pharmacy Program Driving Prescriptions
2013
2014
(1)
National Average fill rate calculated by subtracting IMS Monthly Claims national average rejections and reversals from total p

a claim adjudicated (1 rejections reversals) and Pharmacy Pilot fill rate based on total patients contacted by the pharmacy insurance information and fill their prescription (total patients that fill Rx / total patients that are contacted and have insurance (2)

National Average refill rate based on IMS NPA Monthly

Primary Care Orphan Diseases 10 Four US Products in Three Market Segments (1) Pending the closing of the acquisition of Vidara Therapeutics International Ltd. which is expected this summer (2)

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RAYOS is known as LODOTRA outside of the United States (1) Specialty 250 sales reps

PCPs

Ortho surgeons

Podiatrists Six sales reps 40 sales reps

Rheumatologists (2)

Academic medical centers

Family Practice ID and Immunology

11

(1)Singh and Rosen Ramey. J Rheumatol. 1998;51(suppl):8 16. (2) Geis et al. J Rheumatol. 1996;18:11 14 (3)

M.Wolfe, et.al.; Gastrointestinal Toxicity of Nonsteroidal Anti-inflammatory Drugs; NEJM; vol. 340; no. 24; June 1999.

(4)BMC Musculoskeletal Disorders 2006, 7:79 (5)Sturkenboom, et.al.; Aliment Pharmacol Ther 2003; 18:1137-1147 GI intolerance incidence: up to 50% (1)Endoscopic ulcers incidence: 15-46% (2)Leads to 107k hospitalizations and 16.5k deaths per year (3) 76% of MDs do not prescribe concomitant GI therapy (4) 37% of patients non-compliant; increased to 61% by the 3rd prescription (5)Novel, proprietary formulations of two of the most prescribed NSAIDs combined with a GI protectant in a single pill NSAID-INDUCED **GI TOXICITY** POOR PHYSICIAN AND PATIENT COMPLIANCE VIMOVO VIMOVO DUEXIS DUEXIS Naproxen **NSAID** Ibuprofen Esomeprazole magnesium (PPI) GI Protectant Famotidine (H2 antagonist) BID Dosing TID **VIMOVO & DUEXIS** Addressing an Unmet Medical Need

VIMOVO as the *Smarter Naproxen* There is only ~30% TRx overlap of VIMOVO and DUEXIS prescribers (1) VIMOVO Prescribers DUEXIS DUEXIS Prescribers Prescribers Weekly New Rx (k) **Product Positioning** Minimal Overlap with **Existing Targets** Underlying Market Potential leading to limited overlap in existing writers of VIMOVO and DUEXIS The market potential for ibuprofen and naproxen underlying NSAID is large, segmented, and largely untapped VIMOVO and DUEXIS are highly synergistic and meet different patient needs 12 Significant Market Opportunity for both VIMOVO and DUEXIS with Minimal Overlap

Focus on HCPs that need an NSAID, but are also *concerned with* protection (gold-standard protection, etc.)

Focus on underlying Naproxen prescribers

Focus on HCPs that need *best-in*class pain relief and protection (rapid onset, gold standard efficacy, etc.)

Focus on underlying Ibuprofen prescribers DUEXIS as the *Smarter Ibuprofen* (1) Source: Healthcare Analytics (SHA) Prescriber Level Data from June 2013 August 2013

(naproxen/esomeprazole magnesium) Delayed-Release Tablets 375/20 and 500/20 mg Indicated for the relief of signs and symptoms of osteoarthritis, rheumatoid arthritis and ankylosing spondylitis and to decrease the risk of developing gastric ulcers in patients at risk of developing NSAID-associated gastric ulcers See full prescribing information at www.vimovo.com

Highly Synergistic VIMOVO Acquisition (1) AstraZeneca Annual Reports Product Highlights Product Highlights

Acquired Nov. 18, 2013 from

AstraZeneca

\$35 million one time payment

Leverages existing commercial infrastructure

Focus on commercial payors

Maximizing value through price and lower patient co-pay

Rapid growth in VIMOVO revenues 14 Net Sales Net Sales (1) (1) Perfect example of value arbitrage we are trying to capture in our BD strategy

VIMOVO Off to Strong Start in 2014 LARGE MARKET OPPORTUNITY COMMERCIAL DYNAMICS EXECUTION Large NSAID market (>100M TRx/year)

Naproxen NSAID in U.S. with over 16M TRx/year

Peak annual VIMOVO demand of ~600k scripts and run rate of ~300k scripts at YE13

Branded NSAIDs in Tier 3 position

VIMOVO priced at monthly WAC of \$799, WAC/TRx of ~\$820

84% of claims approved

\$0 target co-pay

May 2014 NRx +2% vs. April 2014

May 2014 TRx +4% vs. April 2014

May 2014 TRx dollars of ~\$21.6M

April 2014 TRx dollars of ~\$20.8M 15 250 Primary Care Reps + 40 Specialty Reps Selling VIMOVO HZNP Full Launch of VIMOVO on February 3, 2014 Source: IMS NPA Monthly data; IMS Claims data Commercial Only

For the relief of signs and symptoms of rheumatoid arthritis and osteoarthritis and to decrease the risk of developing upper gastrointestinal ulcers in patients who are taking ibuprofen for those indications See full prescribing information at www.DUEXIS.com

DUEXIS Scripts Continue to Grow 250 Sales Reps Promoting to Primary Care and ORS 17 LARGE MARKET OPPORTUNITY MANAGED CARE

### EXECUTION

Large NSAID market (100M+ TRx/year)

Ibuprofen is leading NSAID in U.S. with over 33M TRx/year

Branded NSAIDs in Tier 3 position

Monthly WAC of \$799, average WAC/Rx of ~\$720

82% of claims approved

\$0 target co-pay

NRx/TRx continue to grow

May 2014 TRx +8% vs. April 2014

May 2014 NRx +6% vs. April 2014

May TRx dollars of ~\$16.8M

April TRx dollars of ~\$15.7M Source: IMS Xponent data; IMS Claims data - Commercial Only

DUEXIS Quarterly Net Sales Growth (\$M) Sales Force Expansion TRx NRx Price Increase from \$198 WAC to \$502 WAC Price Increase from \$502 WAC to \$677 WAC Price Increase from \$677 WAC to \$799 WAC 18 **DUEXIS Monthly** TRx and NRx (2) (1) DUEXIS Scripts Continue to Grow (continued) DUEXIS Scripts Continue to Grow (continued) Source: IMS Xponent Data (1) Includes one-time amount of \$1.4M due to change in timing of revenue recognition. (2)Includes one time reversal of managed care rebate in the amount of \$2.4M

For reduction of the frequency and severity of serious infections associated with Chronic Granulomatous Disease and for delaying time to disease progression in patients with severe, malignant osteopetrosis (Interferon gamma-1b) Injection Pending the closing of the acquisition of Vidara Therapeutics International Ltd. which is expected this summer See full prescribing information at www.actimmune.com

On March 19, 2014, announced the acquisition of Vidara Therapeutics International Ltd. for 31.35 million shares of Horizon stock, \$200 million in cash and plan to become *Horizon Pharma plc* 

Closing is currently projected to be this summer

#### ACTIMMUNE

Recombinant biologic approved in two ultra orphan indications, CGD and SMO

Realized \$58.9 million in net revenues in 2013

Commercial rights in U.S., Canada, Japan and certain LA, Asian and other ROW territories

Two U.S. patents extending to 2022; perpetual Genentech know-how license

Potential for label expansion, including Friedreich s ataxia and eczema herpeticum

Total headcount of 24, including 6 sales reps with biologic and orphan experience

Horizon Pharma plc corporate structure

Corporate headquarters: Dublin, Ireland

Bermuda headquarters: Hamilton (IP & BLA)

U.S. headquarters: Deerfield, IL 20

Vidara Therapeutics Acquisition Overview

Pending the closing of the acquisition of Vidara Therapeutics International Ltd. which is expected this summer

Pricing Currently priced at \$308K per patient per year Pricing analyses in process to determine optimal pricing strategy

Penetration CGD and SMO penetration is <25%

Opportunity to increase penetration rates with increased commitment to selling and marketing

New Presentations/Formulations ~25% of scripts are for Medicaid covered patients \$0.01 sales price per vial New presentation/formulation reestablish value-based pricing with Medicaid

New Indications 12 patient trial in Freidrich s Ataxia reading out in 2H:14 Eczema herpeticum 21 Growth Potential for ACTIMMUNE Pending the closing of the acquisition of Vidara Therapeutics International Ltd. which is expected this summer

Delayed-Release Low-Dose Prednisone approved in the U.S. for treatment of Rheumatoid Arthritis, Polymyalgia Rheumatica, Psoriatic Arthritis, Ankylosing Spondylitis, Asthma and Chronic Obstructive Pulmonary Disease\* (prednisone) Delayed-Release Tablets Note: RAYOS is known as LODOTRA outside the United States \*For full list of indications, see full prescribing information at: www.RAYOSrx.com

RAYOS Commercial Overview HIGH UNMET NEED IN RA & PMR COMMERCIAL DYNAMICS EXECUTION 1.8M RA Patients, majority suffer from morning symptoms

1.1M PMR Patients, majority suffer from morning symptoms

~10M annual TRx

~3M annual prednisone Rx s

40 Rheum Specialists calling on 3,000+ rheumatologists

May 2014 TRx +4% vs. April 2014

May 2014 NRx -1% vs. April 2014

May TRx dollars of ~\$1.72M

April TRx dollars of ~\$1.67M May 2014 TRx +4% vs. April 2014 Note: RAYOS is known as LODOTRA outside the United States 23 Source: IMS NPA Monthly data; IMS Claims data - Commercial Only

Majority Tier 3 position

RAYOS priced at \$933 WAC per 30-count bottle, WAC per Rx of \$1,600

88% of claims approved

\$0 target co-pay

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DUEXIS

6 issued U.S. patents

Settled Par Pharmaceutical PIV litigation by granting a non-exclusive right to market a generic

product beginning January 1, 2023, or earlier under certain circumstances

#### VIMOVO

8 issued U.S. patents with protection to at least 2022

Five generic companies have filed ANDA PIV against VIMOVO o Dr. Reddy s non-infringement challenge in court system (Pisano, NJ)

Court ordered mediation is in process (no trial date set)

#### RAYOS

0

5 issued U.S. patents with protection to at least 2024

Horizon responded to a PIV Patent Certification received from Watson on July 15, 2013 by filing a patent infringement lawsuit against Watson on Aug. 27, 2013 in New Jersey no trial date set ACTIMMUNE (1)Two U.S. patents extending to 2022; perpetual Genentech know-how license Long Life Proprietary Products (1)

Pending the closing of the acquisition of Vidara Therapeutics International Ltd. which is expected this summer

President & CEO, IDM Pharma (Orphan/Osteosarcoma; sold to Takeda) Todd Smith EVP, Chief Commercial Officer

Agouron, Achillion, Abbott, Bayer, Fenwal VP/GM, Abbott Immunology Led global development & launch of HUMIRA (\$11+B in sales) Wyeth, Searle, Merck and Abbott Marketed 12 NSAIDs, including Celebrex, VIOXX, Arthrotec, Mobic, Daypro, Brufen Jeff Sherman, M.D. EVP, Chief Medical Officer BMS, Searle, Takeda, IDM Pharma 25 Proven Leadership Team Extensive Commercial, Development and M&A Experience **Bob** Carey EVP, Chief Business Officer JMP Securities, Dresdner Kleinwort Wasserstein, Vector Securities Board Member Raptor (orphan); XOMA (orphan) Egalet (opioid); BIO Bob De Vaere (1)EVP, Chief Financial Officer IDM Pharma, Nexa Therapeutics, Epimmune, Vista Medical Paul Hoelscher (1)EVP, Finance Chief Financial Officer Elect

OfficeMax, Alberto Culver/Unilever, KPMG LLC Ben Bove SVP, Marketing & Analytics Galt & Company, Abbott, Fenwal Jeff Kent, M.D. SVP, Medical Affairs Searle (Celebrex/Bextra), Abbott (HUMIRA) Tim Walbert Chairman, President & CEO (1) Bob De Vaere will retire, effective Septe

Bob De Vaere will retire, effective September 30, 2014 and enter into a one-year consulting agreement with the Company; Paresident, Finance, effective June 23, 2014 and Chief Financial Officer, effective October 1, 2014

Leverage Core Commercial Strengths Adjacencies to Current Capabilities Important Unmet Need

Pursue products with differentiated clinical benefits

U.S. products/companies with on-market assets

Leverage 250 person primary care sales force

Leverage 40 person specialty sales force

Build upon orphan presence with ACTIMMUNE

Differentiated and/or underappreciated assets with targeted approach regardless of therapeutic area

Opportunistic targets which provide geographic expansion

Immediate accretion

Attractive financial returns

Long life assets Maximize Shareholder Value Creation Business Development Strategy 26

(\$ in millions) Period 1Q 2014 % Change from 1Q 2013 2Q 2013 3Q 2013 4Q 2013

FY 2013 1Q 2014 1Q 2013 4Q 2013 Net Sales by Product VIMOVO	
- \$	
- \$	
\$ 1.0 \$ 1.0 \$ 34.0 \$ NM NM DUEXIS 4.8	
9.5	
21.6	
23.1	
59.0	
13.9	
190% -40% RAYOS (1) 0.4	
0.4	
1.8	
3.2	
5.8	
3.3	
727%	

3% LODOTRA (1) 3.5			
1.2			
0.7			
2.8			
8.2			
0.7			
-80% -75% Total Net Sales 8.7 \$ 11.1 \$ 24.1 \$ 30.1 \$ 74.0 \$ 51.9 \$ 497% 73% Adjusted EBITDA (3) (16.6) \$ (13.9) \$ (0.8) \$ (1.0) \$ (32.3) \$ NM NM NM 1Q 2014 and Full Year 2013	3 Results		

Adjusted 1Q 2014 non-GAAP net income was \$11.0 million, or \$0.16 non-GAAP basic earnings per share and \$0.13 non-GAAP diluted earnings per share (1) RAYOS is known as LODOTRA outside the United States (2) Not meaningful (3) Adjusted EBITDA reflects adjustments for Vidara acquisition costs 27 (2)

Adjusted Financials Reconciliation 28 (\$ in thousands) Fiscal Year Ended December 31, Three Months Ended March 31, 2011 2012

2013 2013 2014 **GAAP** Net Loss (113,265) \$ (87,794) \$ (149,005) \$ (22, 171)\$ (206,250) \$ Loss on derivative revaluation \_ -69,300 -204,030 Intangible impairment charge 69,621 \_ \_ \_ \_ Depreciation and intangible amortization expense 4,199 5,538 9,310 1,922 5,403 Interest expense, net 6,284

EUgai Filling. HONIZON FHANINA, INC FUITI DEFAT4A
14,525
39,178
3,603
4,207
Other expense, net
56
-
-
667
Foreign exchange loss (gain) 1,023
(489)
(1,206)
905
38
Benefit for income taxes (14,683)
(5,171)
(1,121)
(881)
(1,105)
Non-GAAP Adjustments 66,444 \$ 14,459 \$ 115,461 \$ 5,549 \$

212.240
213,240 \$
EBITDA
(46,821)
\$
(73,335) \$
(33,544)
\$
(16,622)
\$ 6,990
\$
Adjustments for Vidara acquisition costs
-
-
1,252
-,
-
4.040
4,049
Total Non-GAAP Adjustments
66,444
14.450
14,459
116,713
5,549
217.200
217,289
Adjusted EBITDA
(46,821)
\$
(73,335)
\$ (32,292)
\$ \$
(16,622)
\$
11,039
\$

Adjusted Financials Reconciliation (*continued*) 29 Three Months Ended (\$ in thousands) March 31, 2014 GAAP Net Loss (206,250) \$ Loss on derivative revaluation 204,030

Intangible amortization expense (net of tax effect) 4,680

Stock based compensation 1,927

Amortization of debt discount and deferred financing costs 2,333

Depreciation expense 376

Amortization of deferred revenue (161)

Non-GAAP Adjustments 213,185

Non-GAAP Net Income (Loss) 6,935 \$ Adjustments for Vidara acquisition costs 4,049

Total Non-GAAP Adjustments 217,234

Adjusted Non-GAAP Net Income (Loss) 10,984 \$ GAAP Net Loss per common share - basic (3.07) \$ Non-GAAP Adjustments 3.23

Adjusted Non-GAAP Basic Earnings (Loss) per Share 0.16 \$ Dilutive earnings per share effect of common stock equivalents (0.03)

Adjusted Non-GAAP Net Income (Loss) per Common Share - Diluted 0.13 \$

(in millions) As of 3/31/14 Pro Forma for Vidara Acquisition Balance Sheet Balance Sheet Cash

\$103.4
n/a
Debt
\$150.0
(1)
\$450.0
(2)
Capitalization
Capitalization
(3)
(3)
Basic Shares Outstanding
71.4
102.8
Fully Diluted Shares Outstanding
(4)
90.4
121.8
Financial Highlights
30
(1)
Gross amount of 5% convertible notes outstanding, excluding debt discount
(2)
Includes \$300 million senior secured credit agreement; see following slide for further details.
(3)
Includes all issued and outstanding securities, vested and unissued RSUs and contingent stock options. Pro Forma column includes and unissued RSUs and contingent stock options.
shares issued to Vidara shareholders upon closing and assumes no existing warrants, options or RSUs are exercised between 3/
(4)

(4) Excludes shares issuable upon conversion of \$150 million convertible note.

On June 19, 2014, we announced an agreement with a group of lenders to provide Horizon with \$300 million in financing that will replace the \$250 million bridge loan commitment received from Deerfield Management Company, L.P., announced on March 19, 2014 31 Senior Secured Credit Agreement Loan Commitment Senior secured term loans Size \$300 million Term 5 years Interest Rate L + 8.0% (LIBOR Floor: 1.0%) or the prime lending rate + 7.0% (at each borrower s option) Make-whole T+50 bps for two years and callable thereafter at a premium: Year 3: 104, year 4: 102, year 5: 100 Amortization None Accordion (1)Unlimited subject to (i) minimum EBITDA of \$70 million, (ii) maximum senior secured net leverage ratio < 3.5x (cash netting cap of \$50 million) and (iii) maximum total net leverage ratio < 5.5x (cash netting cap of \$50 million) Timing Coincident with the closing of the proposed acquisition of Vidara Use of proceeds To effect the proposed acquisition of Vidara, pay related transaction fees and expenses and for general corporate purposes

(1)

EBITDA to reflect certain adjustments and be calculated on a Last Quarter Annualized basis for any test period ending on or p after which EBITDA shall be calculated on a Last Twelve Months Basis after giving pro forma effect to any acquisitions and c occurred during the test period and on or before the calculation date.

Appendix Appendix

Horizon Pharma History
33
2005
2008
2010
2011
2012

2013 Founded in Palo Alto, CA Relocates to IL DUEXIS U.S. Approval 4-2011 \$50M IPO (NASDAQ: HZNP) DUEXIS U.S. Launch \$111M Raised: \$60M in Debt and \$51M in Equity \$86M Equity Raise RAYOS U.S. Approval 7-2012 RAYOS U.S. Launch T. Walbert joins as CEO Acquisition of VIMOVO \$150M Convert VIMOVO HZNP U.S. Launch 2014 Vidara Acquisition (1) & \$300M Loan (1)Pending the closing of the acquisition of Vidara Therapeutics International Ltd. which is expected this summer Acquisition of private, Switzerland-based Nitec Pharma (RAYOS)

VIMOVO: Significant Reduction in Gastric Ulcers Cumulative observed incidence of Gastric Ulcers \*P<0.001 Ec Naproxen vs. VIMOVO Source: VIMOVO Approved Package Insert, October 2012 \*

34

4.1 23.1 7.1 24.3 0 5 10 15 20 25 30 VIMOVO EC Naproxen VIMOVO EC Naproxen Study 301 Study 302

VIMOVO: Gastric Protection with or without Low Dose Aspirin (LDA) Pooled Cumulative incidence of Gastric Ulcer with or without LDA 35 Aliment Pharmacol Ther 2010; 32: 401-413 LDA Users LDA Non-Users

36 Source: DUEXIS Approved Package Insert, April 2011 20.0% DUEXIS TID Ibuprofen 800 mg TID

10.5% N=190 N=380 p-value = 0.0028.7% N=216 N=447 17.6% DUEXIS TID Ibuprofen 800 mg TID p-value = 0.0004Statistically significant less dyspepsia vs. ibuprofen 5% VS. 8% (p-value = 0.009)All other treatment-emergent GI adverse events were similar

## ~50 % Reduction in Gastric or Upper GI Ulcers

REDUCE-1, Patients with Endoscopic *Gastric* Ulcer (%) REDUCE-2, Patients with Endoscopic *Upper GI* Ulcer (%) DUEXIS Met Primary Endpoints in Phase 3 Trials

RELEASE 10pm 2am 6am 10am Pain & Stiffness High Cytokine release RELEASE High 10pm 2am 6am 10am **RAYOS Synchronizes Pharmaceutical Delivery with** Therapeutic Need 37 Notes: Illustrative only STANDARD PREDNISONE Current regimen too late Morning administration does not mediate nocturnal cytokine peak LODOTRA Optimal nocturnal release regimen Convenient bedtime dosing Reduces morning stiffness and pain Pain & Stiffness Cytokine release DOSING RAYOS is known as LODOTRA outside the United States

Note: RAYOS is known as LODOTRA outside the United States Diff. (1.7X) Diff. (2.5X) Diff.

(2.8X) CAPRA-2 -Pivotal U.S. Phase 3 Study 38 LODOTRA Placebo p-value = 0.0002p-value = 0.0027p-value = 0.0955% of Patients with Improvement Source: Arthritis Rheum 2010 (62 suppl 10:392) 48.5% 22.7% 7.0% 28.6% 9.2% 2.5% ACR 20 ACR 50 ACR 70 Strong, Significant Improvement in ACR 20 and ACR 50 350 Patients Randomized 2:1 DMARD Use (MTX) Included in Trial Safety Comparable to Placebo RAYOS -Significantly Improved ACR 20/50 Response

RAYOS Delivers Superior to Immediate-Release
Prednisone in Reducing Morning Stiffness
39
CAPRA-1 (Pivotal EU Phase 3 Study)
23% mean relative reduction in morning stiffness after 3 months
Sustained reduction of morning stiffness (~50% reduction)

Reduction in IL-6 levels (~30% after 3 months, ~40% after 12 months)

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Relative change from baseline in duration of morning stiffness for the safety set of the open phase **IR PREDNISONE GROUP** SWITCHED TO LODOTRA DOUBLE BLIND PHASE p-value = 0.045Mean Immediate Release Prednisone Group Mean LODOTRA Group Note: RAYOS is known as LODOTRA outside the United States -80 -70 -60 -50 -40 -30 -20 -10 0 10 20 0 1 2 3 4 5 6 7 8 9 10 11 12 13 Study Month OPEN LABEL PHASE Source: The Lancet, 2008 (371:205-14)

NASDAQ: HZNP June 2014 Horizon Pharma, Inc. Filed under Rule 14a-12 of the Securities Exchange Act of 1934 Filing by: Horizon Pharma, Inc. Subject Company: Horizon Pharma, Inc. SEC File No. of Horizon Pharma, Inc.: 001-35238

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described therein that was made available beginning on June 23, 2014. The following is a slide presentation relating to the proposed transactions