UNITED STATES SECURITIES AND EXCHANGE COMMISSION Washington, D.C. 20549

FORM 8-K

CURRENT REPORT

Pursuant to Section 13 or 15(d) of the Securities Exchange Act of 1934

Date of Report (Date of Earliest Event Reported): July 12, 2012

PDL BioPharma, Inc.

(Exact name of Company as specified in its charter)

000-19756 (Commission File Number)

Delaware (State or Other Jurisdiction of Incorporation) 94-3023969 (I.R.S. Employer Identification No.)

932 Southwood Boulevard Incline Village, Nevada 89451 (Address of principal executive offices, with zip code)

(775) 832-8500

(Company's telephone number, including area code)

Check the appropriate box below if the Form 8-K filing is intended to simultaneously satisfy the filing obligation of the Company under any of the following provisions:

□ Written communications pursuant to Rule 425 under the Securities Act (17 CFR 230.425)

□ Soliciting material pursuant to Rule 14a-12 under the Exchange Act (17 CFR 240.14a-12)

D Pre-commencement communications pursuant to Rule 14d-2(b) under the Exchange Act (17 CFR 240.14d-2(b))

D Pre-commencement communications pursuant to Rule 13e-4(c) under the Exchange Act (17 CFR 240.13e-4(c))

Item 7.01 Regulation FD Disclosure.

On July 12, 2012, PDL BioPharma, Inc. (the Company) will make a presentation at The JMP Securities Healthcare Conference in New York, New York. A copy of the Company's presentation materials has been posted to the Company's website and is attached hereto as Exhibit 99.1

Limitation of Incorporation by Reference

In accordance with General Instruction B.2. of Form 8-K, this information, including Exhibit 99.1, is furnished pursuant to Item 7.01 and shall not be deemed to be "filed" for the purpose of Section 18 of the Securities Exchange Act of 1934, as amended, or otherwise subject to the liabilities of that section. The information in this Item 7.01 of this Current Report on Form 8-K will not be deemed an admission as to the materiality of any information that is required to be disclosed solely by Regulation FD.

Cautionary Statements

This Current Report on Form 8-K and the presentation include "forward-looking statements" within the meaning of Section 27A of the Securities Act of 1933, as amended, and Section 21E of the Securities Exchange Act of 1934, as amended. Although we believe that the expectations reflected in the forward-looking statements are reasonable, we can give no assurance that such expectations will prove to be correct. Important factors that could impair the Company's royalty assets or business are disclosed in the "Risk Factors" contained in the Company's 2011 Annual Report on Form 10-K. All forward-looking statements are expressly qualified in their entirety by such factors. We do not undertake any duty to update any forward-looking statement except as required by law.

Item 9.01 Financial Statements and Exhibits.

(d) Exhibits.

Exhibit No.	Description
99.1	Presentation

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the Company has duly caused this report to be signed on its behalf by the undersigned hereunto duly authorized.

PDL BIOPHARMA, INC. (Company)

By: /s/ John P. McLaughlin John P. McLaughlin President and Chief Executive Officer

Dated: July 12, 2012

EXHIBIT INDEX

Exhibit No. 99.1

Presentation

Description





Seventh Annual JMP Securities Healthcare Conference

July 12, 2012

Forward Looking Statements

This presentation contains forward-looking statements, including PDL's expectations with respect to its future royalty revenues, expenses, net income, and cash provided by operating activities.

Each of these forward-looking statements involves risks and uncertainties. Actual results may differ materially from those, express or implied, in these forward-looking statements. Factors that may cause differences between current expectations and actual results include, but are not limited to, the following:

- The expected rate of growth in royalty-bearing product sales by PDL's existing licensees;
- The relative mix of royalty-bearing Genentech products manufactured and sold outside the U.S. versus manufactured or sold in the U.S.;
- The ability of PDL's licensees to receive regulatory approvals to market and launch new royalty-bearing products and whether such products, if launched, will be commercially successful;
- Changes in any of the other assumptions on which PDL's projected royalty revenues are based;
- Changes in foreign currency rates;
- Positive or negative results in PDL's attempt to acquire revenue generating assets;
- The outcome of pending litigation or disputes, including PDL's current dispute with Genentech related to ex-U.S. sales of Genentech licensed products; and
- The failure of licensees to comply with existing license agreements, including any failure to pay royalties due.

Other factors that may cause PDL's actual results to differ materially from those expressed or implied in the forward-looking statements in this presentation are discussed in PDL's filings with the SEC, including the "Risk Factors" sections of its annual and quarterly reports filed with the SEC. Copies of PDL's filings with the SEC may be obtained at the "Investors" section of PDL's website at <u>www.pdl.com</u>. PDL expressly disclaims any obligation or undertaking to release publicly any updates or revisions to any forward-looking statements contained herein to reflect any change in PDL's expectations with regard thereto or any change in events, conditions or circumstances on which any such statements are based for any reason, except as required by law, even as new information becomes available or other events occur in the future. All forward-looking statements in this presentation are qualified in their entirety by this cautionary statement.





Key Information

Ticker	PDLI (NASDAQ)
Location	Incline Village, Nevada
Employees	Less than 10
2011 Revenues	\$362 million
2011 Expenses	\$18.3 million
2012 Regular Dividends (Payable Date)	\$0.15 /share paid on March 14 and June 14, and to be paid on September 14 and December 14
2012 Regular Dividends (Record Date)	March 7, June 7, September 7 and December 7
Q1-2012 Cash Position ¹	\$192.5 million
Shares O/S ²	~ 140 million
Average Daily Volume	~ 1.6 million shares
1. As of March 31, 2012; 2. Not fully diluted	



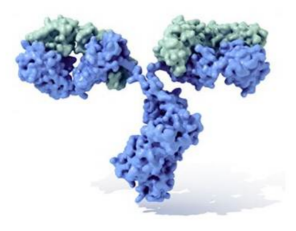




Overview of PDL BioPharma



Antibody Humanization Technology



- Antibodies are naturally produced by humans to fight foreign substances, such as bacteria and viruses
- In the 1980's, scientists began creating antibodies in non-human immune systems, such as those of mice, that could target specific sites on cells to fight various human diseases
- However, mouse derived antibodies are recognized by the human body as foreign substances and may be rejected by the human immune system
- PDL's technology allows for the "humanization" of mouse derived antibodies by moving the important binding regions from the mouse antibody onto a human framework
- PDL's humanization technology is important because the humanized antibodies retain the binding and activity levels from the original mouse antibody
- PDL's technology has been incorporated into antibodies to treat cancer, eye diseases, arthritis, multiple sclerosis and other health conditions with aggregate annual sales of over \$17 billion

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Mission Statement

Queen et al. Patents

- > Manage patent portfolio
- > Manage license agreements

Optimize return for shareholders

Obtain new revenue generating assets

- > Assets that improve shareholder return
- > Backed by commercial stage products
- > Differentiated product profile
- > Indifferent as to therapeutic field
- > Target value of \$75-150 million



Corporate Governance

Management

John McLaughlin President & CEO

Bruce Tomlinson VP & CFO

Christopher Stone VP, General Counsel & Secretary

Caroline Krumel VP of Finance

Danny Hart Deputy General Counsel

Board of Directors

Fred Frank Lead Director

Jody Lindell

John McLaughlin

Paul Sandman

Harold Selick





Licensed Products and Royalty Revenue



Approved Licensed Products: Overview

Product	Licensee	2011 WW Sales	Approved Indications
	Genentech (US) and Roche (ex-US)	\$5.7 billion	Metastatic colorectal cancer Advanced non-small cell lung cancer Renal cancer Metastatic HER2 – breast cancer Glioblastoma Ovarian cancer
Herceptin [®]	Genentech (US) and Roche (ex-US)	\$5.7 billion	Metastatic HER2+ breast cancer Metastatic HER2+ stomach cancer
	Genentech (US) and Novartis (ex-US)	\$3.6 billion	Wet age-related macular degenerative (AMD) Macular edema or swelling following retinal vein occlusion Diabetic macular edema
Colair Omalizumab	Genentech (US) and Novartis (ex-US)	\$1.1 billion	Moderate to sever persistent allergic asthma Fist approved therapy designed to target the antibody IgE, a key underlying cause of the symptoms of allergy related asthma
Tysabri (natalizumab)	Biogen Idec and Elan	\$1.1 billion	Multiple Sclerosis (MS) in adult patients with relapsing forms of the disease Crohn's disease in adult patients with moderate-to-severe forms of the disease who have had an inadequate response to or are unable to tolerate conventional therapies
• ACTEMRA tocilizumab	Roche and Chugai	\$0.7 billion	Rheumatoid arthritis (RA)
PERJETA	Genentech (US) and Roche (ex-US)	Approved on June 8, 2012	Previously untreated HER2+ metastatic breast cancer
percoronab		Roche sales assumes 1.0877	75 CHF/USD

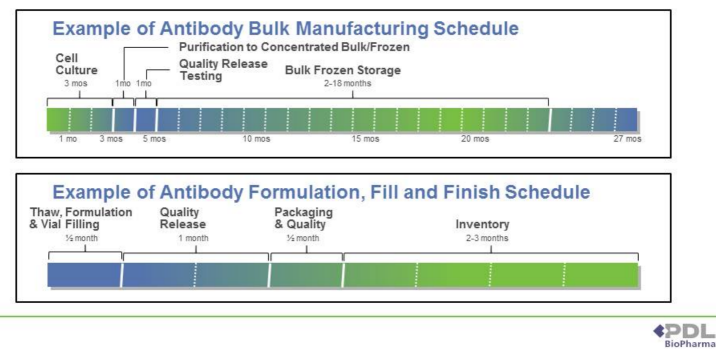
How Long Will PDL Receive Royalties from Queen et al. Patents?

PDL's revenues consist of royalties generated on sales of licensed products

Sold in a patented jurisdiction before the expiration of the Queen et al. patents in mid-2013 through end of 2014

or

Made prior to the expiration of the Queen et al. patents in a patented jurisdiction and sold anytime thereafter



Queen et al Patents - Royalty Rates

Tysabri and Actemra

> Flat, low single-digit royalty

Genentech Products (Avastin, Herceptin, Lucentis¹ and Xolair)

- > Tiered royalties on product made or sold in US
- > Flat, 3% royalty on product made and sold outside US
- > Blended global royalty rate on Genentech Products in 2011 was 1.8%
- > Blended royalty rate on Genentech Products in 2011 made or sold in US was 1.4%

Genentech Product Made or Sold in U.S.	
Net Sales up to \$1.5 Billion	3.0%
Net Sales Between \$1.5 Billion and \$2.5 Billion	2.5%
Net Sales Between \$2.5 Billion and \$4.0 Billion	2.0%
Net Sales Over \$4.0 Billion	1.0%
Genentech Product Made and Sold Ex-U.S.	
Net Sales	3.0%

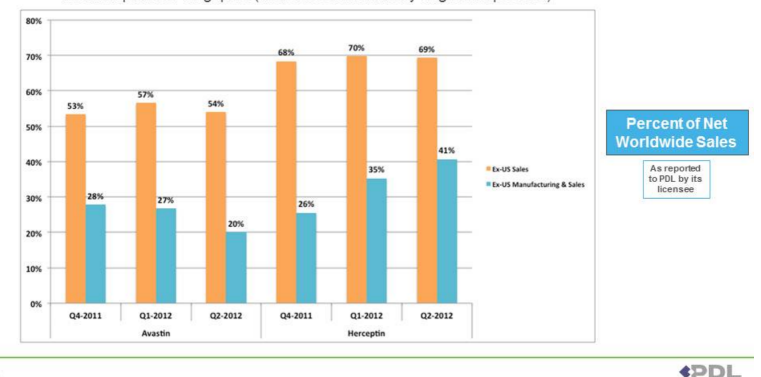
1. As part of a settlement with Novartis, which commercializes Lucentis outside US, PDL agreed to pay to Novartis certain amounts based on net sales of Lucentis made by Novartis during calendar year 2011 and beyond. The amounts to be paid are less than we receive in royalties on such sales and we do not currently expect such amount to materially impact our total annual revenues in 2012.



Ex-US Manufacturing & Sales

Roche is moving some manufacturing ex-US which may result in higher royalties to PDL due to the flat 3% royalty for Genentech Products made and sold ex-US

Current production at Penzburg (Herceptin) and Basel (Avastin) plants
 In June 2011, Roche completed 191 million SFr upgrade and expansion of Penzberg facility



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> Two new plants in Singapore (antibodies and antibody fragments/proteins)

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Royalty Products – Approved

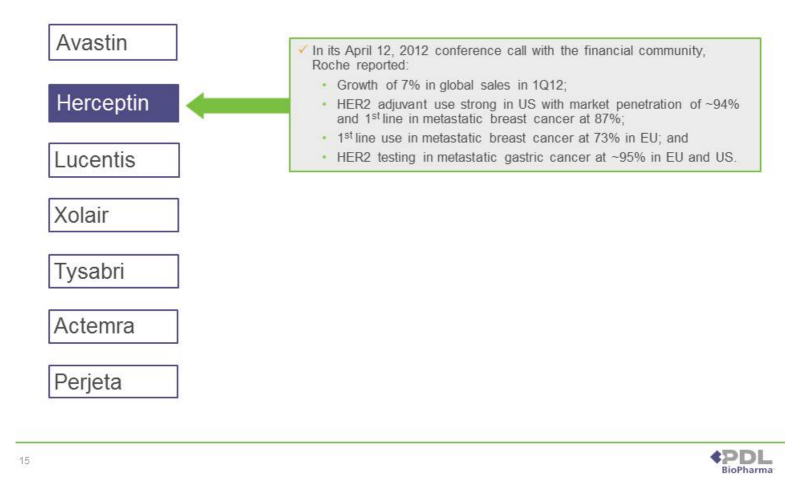


Royalty Products - Avastin

Avastin	 On December 23, 2011, Roche announced that EU approved its use in combination with standard chemotherapy for treatment of newly diagnosed ovarian cancer. On December 28, 2011, Genentech and Roche announced data from Phase 3 trial in patients with advanced, previously untreated ovarian
Herceptin	cancer showed an improvement in primary endpoint of progression-free survival but not secondary endpoint of overall survival.
	On June 2, 2012, Genentech and Roche announced that Phase 3 trial in patients with platinum-resistant, recurrent ovarian cancer of Avastin plus chemotherapy compared to chemotherapy alone:
Lucentis	 Primary endpoint of PFS was 6.7 months compared to 3.4 months; and
Valain	 Secondary endpoint of ORR was 30.9% compared to 12.6%
Xolair Tysabri	In EU where it is approved for ovarian cancer, Roche said that it plans to discuss next steps with health authorities based on this data, while it will await overall survival data from first line setting available in 2013 before deciding whether to file in US.
Actemra	On June 2, 2012, Genentech and Roche reported that Phase 3 trial in patients with metastatic colorectal cancer who received Avastin plus chemotherapy as initial treatment and then Avastin plus a different chemotherapeutic after disease progression met its primary endpoint of overall survival when compared to patients who received only chemotherapy when the disease progressed:
Perjeta	 Overall survival was 11.2 months compared to 9.8 months; and Median PFS was 5.7 months compared to 4.1 months. Genentech and Roche expect to make a global filing in 2012.

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Royalty Products - Herceptin

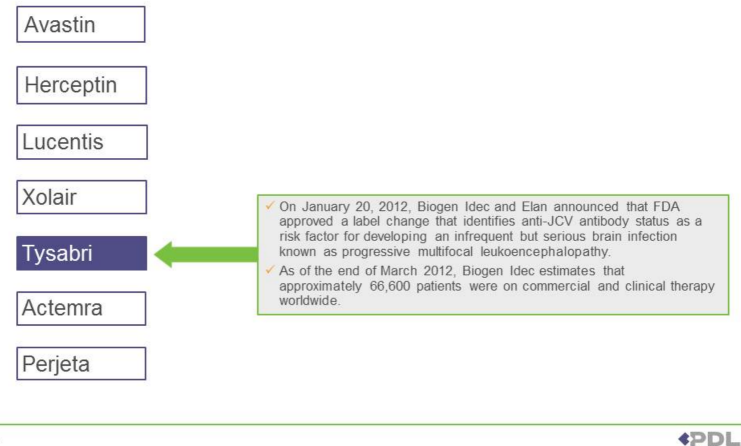


Royalty Products - Lucentis



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Royalty Products - Tysabri



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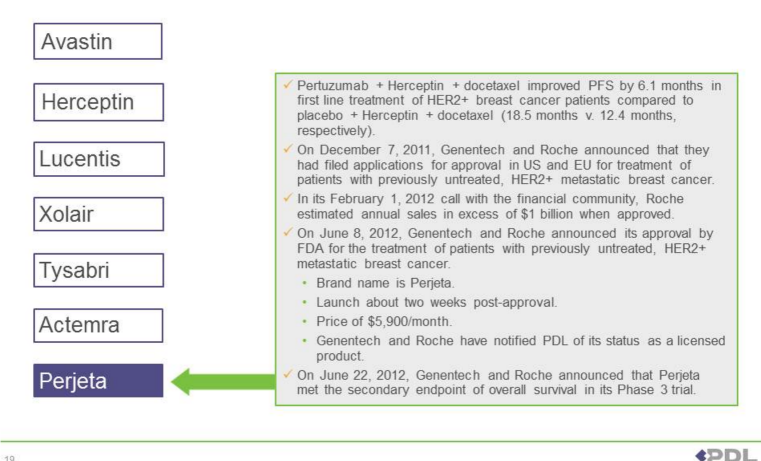
Royalty Products - Actemra



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Royalty Products - Perjeta



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Potential Royalty Products – Development Stage



Potential Royalty Products – T-DM1

T-DM1 Breast HER2+ Cancer

Ocrelizumab Multiple Sclerosis

Afutuzumab Chronic Lymphocytic Leukemia

Bapineuzumab Alzheimer's Disease

Solanezumab Alzheimer's Disease

Datoluzumab Colorectal Cancer

Daclizumab Multiple Sclerosis

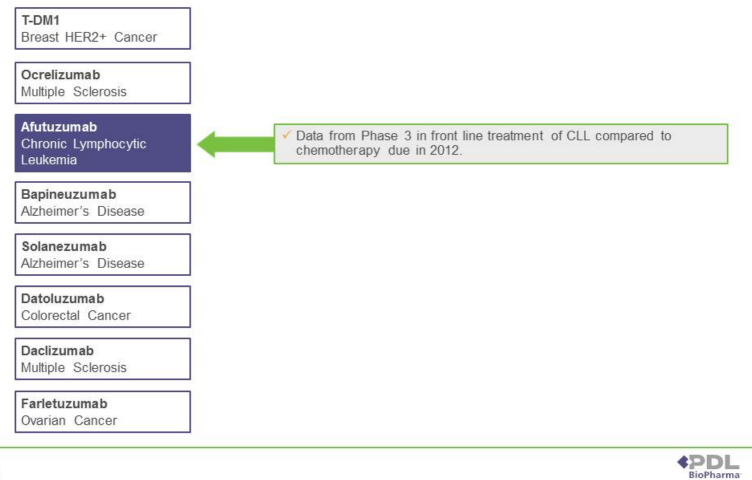
Farletuzumab Ovarian Cancer

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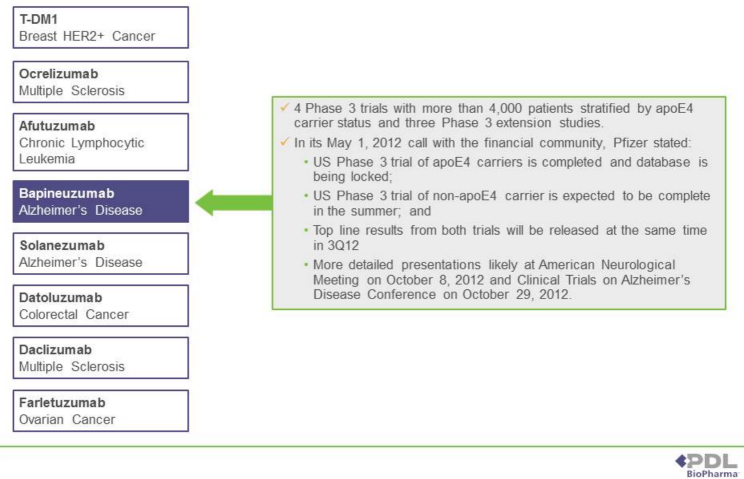
- In its February 1, 2012 call with the financial community, Roche estimated annual sales in excess of \$1 billion when approved.
- On June 2, 2012, Roche/Genentech said that the Phase 3 trial of second line therapy in patients with metastatic HER2+ breast cancer comparing treatment with T-DM1 versus treatment with Tykerb and Xeloda showed:
 - Significant improvement in PFS of 35% (9.6 months v. 6.4 months);
 - One-year survival of 84.7% compared to 77.0%;
 - Response rate of 43.6% compared to 30.8%; and
 - Grade 3 or higher AE's of 40.8% compared to 57.0%
- Roche/Genentech expect to file for second line approval in 2012 and first line in 2014.



Potential Royalty Products – Afutuzumab



Potential Royalty Products – Bapineuzumab



Potential Royalty Products – Solanezumab

T-DM1 Breast HER2+ Cancer	
Ocrelizumab Multiple Sclerosis	
Afutuzumab Chronic Lymphocytic Leukemia	
Bapineuzumab Alzheimer's Disease	 2 Phase 3 trials with approximately 2,000 patients and 1 Phase 3 extension study. In its 2011 earnings call, Lilly reported that an independent
Solanezumab Alzheimer's Disease	monitoring committee conducted interim safety and futility analyses and recommended that the trials continue.
Datoluzumab	Lilly confirmed that Phase 3 data expected in second half of 2012 with detailed presentation likely at the Clinical Trials on Alzheimer's Disease Conference on October 30, 2012.
Colorectal Cancer	PDL receives 12.5 year know-how royalty of 2% from date of first sale in addition to patent royalty of 3%.
Daclizumab Multiple Sclerosis	
Farletuzumab Ovarian Cancer	
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Potential Roles of Herceptin, TDM-1 and Pertuzumab in Breast Cancer

1st Line Metastatic HER2+ Breast Cancer	Herce	eptin + ch	emo		erceptin + umab + ch	emo	TDM-1 + pertuzumab
2nd Line Metastatic HER2+ Breast Cancer	Xe	eloda + laj	patinib			TDM-1	
Adjuvant HER2+ Breast Cancer	Herceptin + chemo		He	rceptin sc	+ Chemo	Herceptin sc + pertuzumab + chemo	
	I	T	1			1	
	2010	2011	2012	2013	2014	2015	2016

Source: Roche investor update, April 12, 2012



Genentech / Roche – Product Pipeline



BioPharma



Financials



First Quarter 2012 Overview

	Quarter Ended March 31				
	((In thousands, except per share			
	2	2012	2011		
Royalty revenues	\$	\$ 77,344		73,336	
G&A Expenses		6,945		5,779	
Operating income		70,399	64	77,557	
Interest expense		(8,700)		(9,154)	
Income before income taxes		61,789	82	68,578	
Income tax expense		21,605		24,033	
Net income		40,184		44,545	
Net income per share - Basic	-	\$0.29		\$0.32	
Net income per share - Diluted	2	\$0.29		\$0.25	
-	March	31, 2012	Dec.	31, 2011	
Cash, cash equivalents and investments		\$192,512		\$227,946	
Total assets	3 .	\$234,963		\$269,471	
Total debt carrying value		\$371,772		\$409,985	

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\$PDL BioPharma



Debt



Current and Long-Term Liabilities

<u>Convertible Notes</u>	<u>Conversion Rate per</u> <u>\$1,000 Principal</u> <u>Amount</u>	<u>Approximate</u> <u>Conversion</u> <u>Price Per</u> Common Share	Effective Date	<u>Principal</u> <u>Balance</u> Outstanding
May 2015 Notes	142.5217	\$7.02	June 5, 2012	\$155,250,000
Series 2012 Notes	162.885	\$6.14	June 5, 2012	\$179,000,000
February 2015 Notes	162.885	\$6.14	June 8, 2012	\$1,000,000
Secured Non-Recourse Notes	N/A	N/A	N/A	\$22,737,726

- Bond hedge effectively increases conversion price in May 2015 Notes to \$8.25.
- In 2011 and 2012, we restructured two convertible notes to "net-share" settled and eliminated 44.8 million dilutive shares from the diluted earnings per share calculation in the first quarter of 2012 when compared to the first quarter of 2011.
- In 3Q12, we expect to retire the Secured Non-Recourse Note returning to PDL approximately 40% of the Genentech royalties currently dedicated to payment of the Notes quarterly principal and interest.

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Legal Matters



Pending Dispute with Genentech and Roche

- In August 2010, Genentech sent a fax on behalf of Roche and Novartis asserting its products do not infringe PDL's supplementary protection certificates (SPCs)
 - > Products include Avastin, Herceptin, Lucentis and Xolair
 - SPCs are patent extensions in Europe that are issued on a country-by-country and product-by-product basis

PDL Response

- > Genentech's assertions are without merit
- > PDL disagrees with Genentech's assertions of non-infringement
- Genentech had waived its rights to challenge our patents, including SPCs in its 2003 Settlement Agreement with PDL

2003 Settlement Agreement

- > Resolved intellectual property disputes between the two companies at that time
- Limits Genentech's ability to challenge infringement of PDL's patent rights, including SPCs, and waives Genentech's right to challenge or assist others in challenging the validity of our patent rights



Nevada Lawsuit Against Genentech/Roche

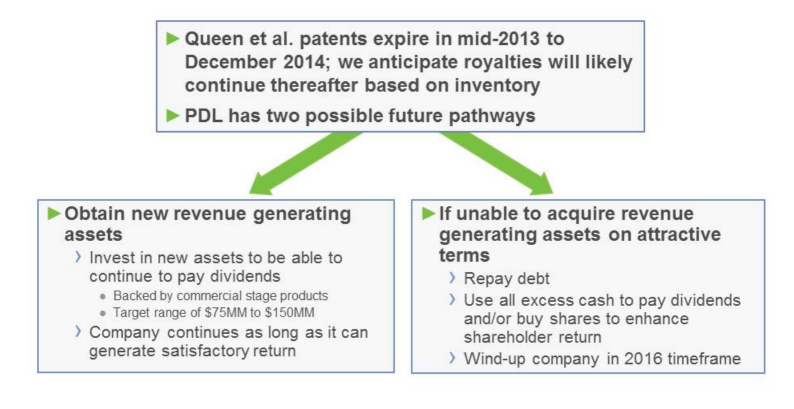
PDL filed a lawsuit against Genentech and Roche in Nevada state court > Lawsuit states that fax constitutes a breach of 2003 Settlement Agreement because Genentech assisted Roche in challenging PDL's patents and SPCs > Complaint seeks compensatory damages, including liquidated damages and other monetary remedies set forth in the 2003 Settlement Agreement, punitive damages and attorney's fees In November 2010, Genentech and Roche filed two motions to dismiss > They contend that 2003 Settlement Agreement applies only to PDL's U.S. patents > They asserted that the Nevada court lacks personal jurisdiction over Roche On July 11, 2011, court denied Genentech and Roche's motion to dismiss four of PDL's five claims for relief and denied Roche's separate motion to dismiss for lack of personal jurisdiction > The court dismissed one of PDL's claims that Genentech committed a bad-faith breach of the covenant of good faith and fair dealing Subsequent to the ruling. Roche has waived its defense that the Nevada court lacks personal jurisdiction for > the purposes of this lawsuit The court ruling allows PDL to continue to pursue its claims that) Genentech is obligated to pay royalties to PDL on international sales of the Genentech Products > Genentech, by challenging, at the behest of Roche and Novartis, whether PDL's SPCs cover the Genentech Products breached its contractual obligations to PDL under the 2003 settlement agreement > Genentech breached the implied covenant of good faith and fair dealing with respect to the 2003 settlement agreement > Roche intentionally and knowingly interfered with PDL's contractual relationship with Genentech in conscious disregard of PDL's rights Parties are currently in discovery and trial is re-scheduled for October 2013 (PDI **BioPharma**



Optimizing Stockholder Return



Business Strategy



4PDL

BioPharma

Investment Highlights

- Strong historic revenue growth from approved products
 Potential for additional indications from existing products, new product approvals and purchase of new revenue generating assets
 Potential to grow and diversify revenues with the addition of new revenue generating assets
 Significantly reduced expenses with no R&D burn
 Liquidity volume averages 1.5 million shares/day
 Return to stockholders

 In 2011, paid regular, quarterly dividends totaling \$0.60/share
 In 2012, paid regular, quarterly dividends of \$0.15/share on
 - March 14 and June 14, and will pay on September 14 and December 14

