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Israel Investors Conference
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Filed by Teva
Pharmaceutical Industries Ltd.
(Commission File No. 001-16174) pursuant to
Rule 425 under the Securities Act of 1933
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Subject Company: Mylan
N.V.

Commission File No.: 333-199861

Safe Harbor Statement

This communication contains forward-looking statements within the meaning of the Private Securities Litigation Reform Act of current beliefs and expectations and involve a number of assumptions, known and unknown risks and uncertainties that change performance or achievements to differ materially from the results, performance or achievements expressed or implied by such assumptions, known and unknown risks and uncertainties include, but are not limited to, those discussed in our Annual Report 2014 and in our other filings with the U.S. Securities and Exchange Commission (the SEC), and those relating to Mylan is filings with the SEC, which factors are incorporated herein by reference. Forward-looking statements are generally identified

believes, intends, estimates, will, would, could, should, may, plans and similar expressions. All state statements that could be deemed to be forward-looking statements, including statements about the proposed acquisition of Myl transaction, the expected future performance (including expected results of operations and financial guidance), and the combin operating results, strategy and plans. Important factors that could cause actual results, performance or achievements to differ r statements we make in this communication include, but are not limited to: the ultimate outcome of any possible transaction bet possibility that no transaction between Teva and Mylan will be effected or that a transaction will be pursued on different terms with the proposed transaction and the results thereof; the effects of the business combination of Teva and Mylan, including the condition, operating results, strategy and plans; uncertainties as to the timing of the transaction; the possibility that the expecte integration of our operations with Mylan s operations (including any expected synergies) will not be fully realized by us or magnetic or the synergies of the s effects on the market price of Teva s or Mylan s shares, including negative effects of this communication or the consummation obtain regulatory approvals on the terms proposed or expected and satisfy other conditions to the offer, including any necessar timely basis; our and Mylan s ability to comply with all covenants in our or its current or future indentures and credit facilities manner, could trigger a default of other obligations under cross default provisions; our and Mylan s exposure to currency fluc the effects of reforms in healthcare regulation and pharmaceutical pricing and reimbursement; uncertainties surrounding the le registration and approval of biotechnology-based medicines; the impact of competition from other market participants; adverse corruption, major hostilities or acts of terrorism on our or Mylan s significant worldwide operations; other risks, uncertainties on Form 20-F for the year ended December 31, 2014 and in our other filings with the SEC; and the risks and uncertainties and documents filed with the SEC. All forward-looking statements attributable to us or any person acting on our behalf are express statement. Readers are cautioned not to place undue reliance on any of these forward-looking statements. Forward-looking sta they are made and we assume no obligation to update or revise any forward-looking statement, whether as a result of new info

Additional Information

This communication is for informational purposes only and does not constitute an offer to buy or solicitation of an offer to sell a proposal which Teva has made for a business combination transaction with Mylan. In furtherance of this proposal and subjectile one or more proxy statements, registration statements or other documents with the SEC. This communication is not a substatement, prospectus or other document Teva and/or Mylan have filed or may file with the SEC in connection with the propose made except by means of a prospectus meeting the requirements of Section 10 of the U.S. Securities Act of 1933, as amend

URGED TO READ THE PROXY STATEMENT(s), REGISTRATION STATEMENT, PROSPECTUS AND OTHER DOCUENTIRETY IF AND WHEN THEY BECOME AVAILABLE AS THEY WILL CONTAIN IMPORTANT INFORMATION statement(s) (if and when available) will be mailed to stockholders. Investors and security holders may obtain free copies of the registration statement, prospectus and other documents (in each case, if and when available) filed with the SEC by Teva through http://www.sec.gov.

No permission has been sought or received to quote from, or refer to, materials cited in this presentation.

Teva, Erez Vigodman, President and Chief Executive Officer and a director of Teva, Eyal Desheh, Group Executive Vice President, Olafsson, President and Chief Executive Officer, Global Generic Medicines of Teva, Kevin C. Mannix, Senior Vice President, directors named in Teva s annual report on Form 20-F filed with the SEC on February 9, 2015 may be deemed participants of Mylan in respect of Mylan s proposal for a business combination with Perrigo Company plc. Additional information may be beneficially owns 22,600,000 ordinary shares of Mylan. To the knowledge of Teva, none of the individuals mentioned above holdings or otherwise, in Mylan or Perrigo or the matters to be acted upon, if any, in connection with a potential business combination of the individuals mentioned above holdings or otherwise, in Mylan or Perrigo or the matters to be acted upon, if any, in connection with a potential business combination or provided the properties of the provided the

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Teva and Mylan

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Drive Organic Growth
Creating
a New
Future
for Teva
Solidify the Foundation
Maintain COPAXONE® Franchise

Generics: Established Global Generic Medicines; improved profitability by ~500 bps

Cost Reduction: Reduced net costs by ~\$600 million

Operations: Accelerated the transformation of our operational network; closed or

divested 11 facilities

Quality: Significant achievements making quality a core competitive competency

Cash Flow: Strong focus resulted in robust cash flow from operations and free cash

flow

Generics:

Solid 2014 performance with stronger profitability

19 product launches in the U.S., 209 in Europe and 87 in ROW delivering ~\$1.0 billion in revenues

Specialty:

Successfully launched four new products with revenues of ~\$200 million

Therapeutic areas selection and focus

Six major submissions of specialty products; eight major approvals

Complemented the specialty pipeline with the acquisitions of Labrys and Auspex

Teva is well on its way to create a new future for the company by targeting a unique space in the industry, building on its strong capabilities in generics and specialty and the intersection between the two

Successfully launched COPAXONE® 40mg in the U.S. and achieved 68.9% conversion rate as of July 1, 2015

Successful and further upcoming launches in various EU countries and elsewhere Significant endeavors on the legal and regulatory front

Successfully

managed

the

life

cycle

of

TREANDA,

ProAir,

and

Azilect

Significant Achievements on All 2014 Must Wins 5

6 2013 2014 % YoY 2015E Q1 2015 % QoQ Revenues \$m \$20,314 \$20,272 \$19.0B-19.4B \$4,982 Operating Income \$m \$5,198 25.6% \$5,732 28.3% +10% \$5.7B-5.9B \$1,533 30.8% +11% **EPS** \$ per share \$5.01 \$5.07 +1% \$5.05-5.35 \$1.36 +11% Cash flow from Operations \$m \$3,237 \$5,127 +58% \$4.3B-4.7B \$1,354 +51% Free Cash Flow \$m \$2,309 \$4,256 +84% \$3.5B-\$3.9B \$1,213

+80% Note: Operating income and EPS are non GAAP results Solidified base shown by robust financial performance in 2014 and Q1 2015 Solidified Base Manifested in Strong Performance in 2014 & 2015

Strong results despite currency head wind

EPS (\$) Specialty Pipeline Existing Specialty Cost Reduction Generics FY 2015 FY 2016

FY 2017 FY 2018 5.00 Profitable Growth in Generics Manage the Life Cycle of Key **Specialty Products** Deliver on the Promise in our Specialty Pipeline **Execute our Cost Reduction** Program 1 2 3 **ILLUSTRATIVE** Note: Earlier entry by generics could reduce operating income by \$30-50M per month In 2014, Teva established a stable base for future organic EPS growth Clear Pathway to **EPS** Growth: Teva s Four Levers of Growth FY 2014 FY 2019

7

Continue to improve operating profitability More focus on key markets and portfolio management Execution of growth market strategy Clear strategy

for OTC

Sales force effectiveness

in key markets

Note: Profitability consists of gross profit, less S&M and R&D expenses related to the segment; segment profitability does not Continued Growth and Improvement in Generics

- 45.2%
- 43.5%
- 41.3%
- 43.3%
- 46%
- 20.2%
- 19.9%
- 16.7%
- 21.9%
- 27%
- 10%
- 20%
- 30%
- 40%
- 50%
- FY11
- FY12
- FY13
- FY14

FY15 mid-point

Gross Profit Margin

Segment Profit Margin

8

9 2014A 2015E 2016E 2014-2016 Cumulative With 2013

Gross Cost Savings (1,000)(650) (400)(2,050)(2,450)Reinvestment in Additional Activities 400 100 200 700 1,600 Net Spend Reduction (600)(500) (250)(1,350)

(850)

Strong Track Record of Driving Cost Savings

10
40mg Success
40 mg 3x a week already at 69% conversion rate; became MS leading therapy in one year post launch
A full launch plan in EU and ROI. Israel
80% conversion
IP Protection

Teva has three Orange Book patents that expire in 2030 (1) The Patent Office has upheld Teva s position on Copaxone® 40mg Teva is well-positioned to respond to IPRs Strengthening Our Specialty Business U.S. Patent Nos. 8,232,250; 8,399,413; and 8,969,302 Maintaining the Copaxone® Franchise Maintaining Other Specialty Products License to commercialize Eagle s Bendamustine Rapid Infusion Product Enhance and protect the TREANDA® (bendamustine hydrochloride) franchise

Expansion of the Azilect franchise to markets outside of the U.S.

FDA Approval of ProAir® RespiClick

(April 2015). Q2 2015 launch

1.
Launches in 2014 include the U.S., Israel, Argentina and Chile 2.
Sales figures exclude U.S. sales of COPAXONE 40mg
Q1 2015
Q2 2015
Q3 2015

Q4 2015
Select
European
markets,
Mexico,
Turkey
and
Australia
(1)
Hydrocodone
ER AD
2014
Multiple Specialty Product Launches in 2014 and 2015
Hydrocodone
ER AD

40 mg/ml

11

Cumulative estimated sales from new specialty product launches of \sim \$200 million in 2014 and \sim \$600 million in 2015

(2)

12 Ca

Capitalizing on a Deep and Promising Pipeline

Phase I

Phase II

Phase III

Registration

TV-46763 (abuse deterrent)

Pain

TV-46139 (abuse deterrent)

Pain

TEV-48125 (anti CGRP)

Chronic and episodic migraine

Note: Pipeline as of April 15, 2015. Phase I includes also projects designated for IND filing

1.

Filed by Eagle Pharmaceutical, commercialized by Teva

Migraine & Pain

CEP-33237 ER Hydrocodone

(abuse det.) U.S. -

Pain

Zecuity

US-

Migraine

13 Analyst commentary following successful phase IIb outcome for TEV-48125 in Chronic and Episodic Migraine

The anti-CGRP class could be VERY Large: \$8-10B.

Clinical benefit is very meaningful.

Safety data very good thus far

We see Teva

taking 30% of the Worldwide CGRP market with peak sales of \$3B

TEVA s anti-CGRP looking better than competitors on primary endpoint ISI Evercore, June 22, 2015

TEV- 48125 leading the pack, with a compelling profile in both chronic and episodic migraine Citi, June 21, 2015

It s still early and far from approval, but we think TEVA might have the drug to beat at this point $\,$. The Teva

data presented this weekend further convinces us that, if approved, Teva could have a

potential blockbuster drug on its hands given the clinical profile and patient need BMO Capital Markets, June 22, 2015

Turning to the stock, we would argue, that this product, coupled with some other products on the branded side and Teva s improvement in the generic business, has not been captured in Teva s share price as the stock has been frozen by the Teva-Mylan-Perrigo love triangle.

The stock is significantly under-valued on our analysis . Bernstein, May 20, 2015

Note: Pipeline as of April 15, 2015. Phase I includes also projects designated for IND filing

Filed by Eagle Pharmaceutical, commercialized by Teva

Capitalizing on a Deep and Promising Pipeline

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Phase III

Registration TV-46763 (abuse deterrent) Pain TV-46139 (abuse deterrent) Pain SD-809 Tourette Syndrome SD-560 Idiopathic pulmonary fibrosis/other fibrotic conditions TEV-48125 (anti CGRP) Chronic and episodic migraine Laquinimod Multiple sclerosis (relapsing remitting) SD-809 Tardive dyskinesia Note: Pipeline as of April 15, 2015. Phase I includes also projects designated for IND filing Filed by Eagle Pharmaceutical, commercialized by Teva Laquinimod Multiple sclerosis (progressive forms) Laquinimod Huntington s Disease Pridopidine Huntington s Disease SD-809 HD (Mid-2015 NDA filing) CEP-33237 ER Hydrocodone (abuse det.) U.S. -Pain Migraine & Pain Movement Disorders & Neurodegeneration COPAXONE 40mg 3w ROW Multiple sclerosis COPAXONE 20mg per Day Japan

Zecuity US-Migraine

Multiple sclerosis

Huntington s
Disease
SD-809: Significant Near-Term Commercial Opportunity
Tardive
Dyskinesia
Estimated Patient
Population (U.S.)

- ~30,000 Patients
- ~500,000 Patients
- ~150,000 Patients

Other Considerations

Only one approved drug in the U.S.: Tetrabenazine

Only 5% of patients treated

2014 sales of ~\$300m million

Annual price per patient of \$80-\$85k

Established reimbursement landscape Received FDA orphan designation Expected launch in 2016 No approved treatment in the U.S.

Tetrabenazine is approved in the EU Limited off-label usage of Tetrabenazine in the U.S. despite significant clinical response

Improved profile should result in increased usage Only one approved product in the U.S.: Aripiprazole

Associated with drowsiness, agitation, weight gain, and sleep

disturbances

Limited

off-label

usage

of

Tetrabenazine

despite

significant

clinical

response

Received FDA orphan designation

Tourette s

Syndrome

15

SD-809 is expected to contribute up to \$800 million to Teva

by 2019, and an estimated

\$2 billion five years following the Tardive

Dyskinesia

launch

Analyst commentary following successful phase IIb outcome of SD-809 in HD, TD

The positive headline data for SD-809 in tardive dyskinesia (obtained via TEVA s recent acquisition of Auspex) provide further validation of TEVA s emerging pipeline

Citi, June 16, 2015

We believe the positive study results are encouraging $\,$. The positive data should help lower the risk profile for this program, which addresses a large and unmet medical need. BMO Capital Markets, June 16, 2015

Auspex brings to Teva

a deep pipeline and with proven deuterium chemistry technology which supports multiple platforms for growth. SD-809 is currently in Phase 3 for tardive dyskinesia and Phase 1 for Tourette syndrome. SD-560 (deuterated pirfenidone) is currently in

development for idiopathic pulmonary fibrosis.

We believe Auspex enhances Teva's mid to long-term revenue and earnings growth, profitability, and product diversity. It is expected to be accretive to non-GAAP EPS beginning in 2017 and meaningfully accretive thereafter, and diversifies Teva's Specialty pharma products offerings .

Maxim, June 16, 2015

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Similar to Huntington $\,$ s disease, TD represents an area of significant unmet need with limited treatment options (there are no FDA-approved drugs for TD). Our estimates and target include value for HD but not the larger TD indication. For reference, a scenario with TD sales reaching \$2bn (TEVA $\,$ s peak sales estimate) would add \sim \$10/share to the theoretical DCF value for TEVA, all else equal $\,$.

Deutsche Bank, June 16, 2015

Analyst commentary following successful phase IIb

outcome of SD-809 in HD, TD 17

Note: Pipeline as of April 15, 2015. Phase I includes also projects designated for IND filing

Filed by Eagle Pharmaceutical, commercialized by Teva

Capitalizing on a Deep and Promising Pipeline

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Phase III

Registration TV-46763 (abuse deterrent) Pain TV-46139 (abuse deterrent) Pain SD-809 Tourette Syndrome SD-560 Idiopathic pulmonary fibrosis/other fibrotic conditions TEV-48125 (anti CGRP) Chronic and episodic migraine SD-809 Tardive dyskinesia COPAXONE 40mg 3w ROW Multiple sclerosis COPAXONE 20mg per Day Japan Multiple sclerosis Reslizumab IV Asthma Note: Pipeline as of April 15, 2015. Phase I includes also projects designated for IND filing Filed by Eagle Pharmaceutical, commercialized by Teva Laquinimod Multiple sclerosis (progressive forms) Laquinimod Huntington s Disease Pridopidine Huntington s Disease Fluticasone Salmeterol **Spiromax** EU Asthma, COPD Reslizumab SC Asthma Fluticasone Salmeterol (MDI) EU Asthma, COPD TEV-46017 (tidal inhaler) **COPD** TEV-48108 (tidal inhaler) **COPD** Laquinimod Multiple sclerosis (relapsing

remitting)

Asthma

Fluticasone Propionate MDPI

Fluticasone Salmeterol

MDPI

Asthma

QVAR (BAI) U.S.

Asthma

SD-809

HD (Mid-2015

NDA filing)

CEP-33237 ER Hydrocodone

(abuse det.) U.S. -

Pain

Respiratory

Migraine & Pain

Movement Disorders & Neurodegeneration

Zecuity

US-

Migraine

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Note: Pipeline as of April 15, 2015. Phase I includes also projects designated for IND filing

Filed by Eagle Pharmaceutical, commercialized by Teva

Capitalizing on a Deep and Promising Pipeline

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Phase III

Registration TV-46763 (abuse deterrent) Pain TV-46139 (abuse deterrent) Pain SD-809 Tourette Syndrome SD-560 Idiopathic pulmonary fibrosis/other fibrotic conditions SD-809 Tardive dyskinesia COPAXONE 40mg 3w ROW Multiple sclerosis COPAXONE 20mg per Day Japan Multiple sclerosis Reslizumab IV Asthma Bendamustine Rapid Infusion (1)CLL, NHL Note: Pipeline as of April 15, 2015. Phase I includes also projects designated for IND filing 1. Filed by Eagle Pharmaceutical, commercialized by Teva Fluticasone Salmeterol Spiromax EU Asthma, COPD Reslizumab SC Asthma Fluticasone Salmeterol (MDI) EU Asthma, COPD TEV-46017 (tidal inhaler) **COPD** TEV-48108 (tidal inhaler) **COPD** Laquinimod Multiple sclerosis (relapsing remitting) Fluticasone Propionate MDPI Asthma Fluticasone Salmeterol **MDPI**

Asthma

Asthma

QVAR (BAI) U.S.

CEP-33237 ER Hydrocodone

(abuse det.) U.S. -

Pain

SD-809

HD (Mid-2015

NDA filing)

TEV-48125 (anti CGRP)

Chronic and episodic migraine

Laquinimod

Multiple sclerosis (progressive

forms)

Laquinimod

Huntington s Disease

Pridopidine

Huntington s Disease

Zecuity

US-

Migraine

19

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Strengthening our specialty business

In 2019 we expect to generate \$5.3 billion in incremental annual risk-adjusted revenues from new specialty product launches (excluding Copxone), that have started in 2014

0.0

5.0

10.0

2014

2015

2016

2017

2018

2019

LOEs*

New Launches

Net Sales

* Copaxone

family included in the LOEs

\$B

20

21 Teva 2020 Generics Specialty New Networked

DOD	3 4	. 1	1 1
ν_{x_7}	1 1	α	ΔI
R&D	171	.vu	LUI

Products &

TA business

Model

Operations

& Quality

Diagnosis,

Prediction,

Prevention

Deploying

Big Data

Consumer/

Patient Driven

Company

Markets

Innovation

Teva s business model transformation

Targeting
a Unique
Space In The
Industry
Generics
Specialty

Our key priorities for business development in 2015 Attractive Pipeline Assets/ **Portfolios** In-Market or Close to Market Assets in Core TAs Unique Health Solutions, Technologies, Services Growth Markets Complex/Hard to Produce Assets or Technologies

Large transactions, where actionable and generate significant strategic and financial long-term value

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Mylan Acquisition Clear and Compelling Strategic
Rationale
Clear and compelling strategic and financial rationale supported by significant shortand long-term value creation to stakeholders of both companies
Industry-leading

company, well-positioned to transform the global generics space

Significantly expanded and more efficient global footprint, including leadership positions and strengthened operations, sales and R&D platforms in attractive markets around the world

Benefits from a robust, industry-leading sales infrastructure and deep customer and provider relationships across the expanded network

Enhanced financial profile

The combined company is expected to have substantial debt capacity and an investment grade rating Strong cash flow generation will allow deleveraging to at or below 3.0x gross debt to EBITDA after 24 months Strongly positioned from day one to pursue future acquisitions to expand portfolio in both specialty pharmaceuticals and generics

Establishes a unique and differentiated business model, leveraging its

significant assets and capabilities in generics and specialty

Leading positions in multiple sclerosis, respiratory, pain, migraine, movement disorders and allergy therapeutics

Enhanced global infrastructure to pursue current and future commercialization

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Transaction Overview

Mylan
Proposed Transaction Overview
\$82.00 per share
Approximately 50% cash / 50% stock
Implies a total equity value of approximately \$43 billion
Teva has already spent \$1.6 billion to establish a 4.6% ownership interest in Mylan

Proposed Price and

Consideration

Significant Premium

48.3% premium to the unaffected Mylan stock price of \$55.31 on March 10, 2015, after which there was widespread speculation of a transaction between Teva and Mylan

Clear Roadmap to

Completion

Have carefully studied the regulatory aspects of proposed combination

Confident that any necessary regulatory requirements will be met in a timely manner; divestitures can be determined and implemented promptly

Filed for HSR on April 21, 2015; initiated pre-merger notification process with European Commission on April 24, 2015

Can be completed in 2015

Financing and

Conditions

No financing condition

Contingent on Mylan not completing its proposed acquisition of Perrigo or any alternative transactions Does not require a Teva stockholder vote

Value Creation

Transaction expected to deliver approximately \$2 billion annually in cost synergies and tax savings, to be largely achieved by the third anniversary of the closing of the transaction

Significant savings from operational, SG&A, manufacturing and R&D efficiencies

Expected non-GAAP EPS accretion in the mid-teens in the first year, and approaching 30% by the third year

26
Meaningful and Real Commitment by Teva
Established a meaningful 4.6%
(~\$1.6 billion) stake in Mylan
Progressed antitrust process
Teva is fully committed to completing the acquisition of Mylan, and has taken significant steps on many fronts in order to do so

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Teva and Mylan s Businesses are Highly Complementary

Teva

(1) Mylan

(2)

Business units: generics, specialty

Specialty therapeutic areas: respiratory / allergy Operates in 145 markets 30,000 employees 2014 revenue: \$9.7 billion Current rating: Baa3 / BBB-Generics 85% Specialty 13% OTC / Other 2% Business units: generics, specialty, OTC Specialty therapeutic areas: CNS, pain, respiratory Operates in 100 markets 43,000 employees 2014 revenue: \$20.3 billion Current rating: A3 / A-Generics 49% Specialty 42% Other / OTC 9% Source of Mylan information: Mylan filings Based on 2014 results Pro forma for the acquisition of Abbott s Non-U.S. Developed Markets Specialty and Branded Generics Business; revenue an North America 48% Europe 33% **ROW** 19% U.S. 52% Europe 29% **ROW** 19% Product offerings are highly complementary and would further enhance the broadest portfolio in the industry

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The Strength of the Combined Company

Source of Mylan information: Mylan filings; financials include contributions from Abbott assets

1.

Net of one-time restructuring costs

2.

Pro Forma for Abbott Non-U.S. Developed Markets Specialty and Branded Generics Business based on 2014 financials

Edgar Filing: Mylan N.V. - Form 425 The combined company is an attractive investment opportunity: financially, strategically and as a platform for future M&A Long-Term Impact **Combined Company** Revenue **EBITDA** >\$30 billion >\$6 billion Significantly expanded and more efficient global footprint Pro Forma 2014 Revenue Mix Expected investment grade rating Opportunity for rapid deleveraging and the funding of future growth Opportunities for capital expenditures synergies of approximately \$350 million annually Enhances product diversification Enhances geographic diversification More diversified organization with the scale and resources to drive value North America 51% Europe 30% Rest of World 19% By Product Type (2) By Geography (2) >\$10 billion Opportunities for substantial achievable cost synergies and tax savings are estimated to be approximately \$2 billion annually 2016E 2018E ~\$33 billion >\$8.5 billion ~\$13 billion Generics 60% Specialty 33%

OTC / Other

Cash Flow from Operations

7%

(1) Free Cash

Flow

(1)

>\$5 billion

>\$7.5 billion

EBITDA Margin

~34%

~40%

30
Recent Industry Trends Support a Combination
Increasingly Fragmented Generics Market
Recent Channel Consolidation
2009
2013
Market Share of the Top 3 U.S. Generics Players

Source: IMS Health; market share as measured by sales 1. Pharmacy benefit managers typically third party administrators prescription drug programs; primarily responsible for processing and paying prescription drug claims 2. Top three include ABC-Walgreens, Cardinal-CVS and McKesson-RiteAid Top three include Celesio, Alliance Boots and Phoenix Top 3 35% Other 65% Top 3 43% Other 57% 2007 Wholesalers Retailers **PBMs** (1) Key Global Distributors Today Wholesalers Retailers **PBMs** The market share of Teva s top three customers increased significantly from 2009 to 2013, with top 3 customer share growing from 52% to 83% in the U.S. (2)

and 51% to 60% in the EU (3)

Significant Premium to Current and Historic Valuation 48.3% premium to the unaffected Mylan stock price of \$55.31 on March 10, 2015, after which there was widespread speculation of

a

transaction

between Teva and Mylan (1)

Proposed Price per Share: \$82.00

\$ per share 3/10/15

48.3% Premium Source: FactSet as of [July 1, 2015]

Mylan LTM Price Performance

\$82.00 per share represents a significant premium for Mylan stockholders

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Prior to speculation regarding Teva s acquisition of

Mylan (March 10,

2015) \$20

\$30

\$40 \$50

\$60

\$70

\$80

Jul 2014

Sep 2014

Nov 2014

Feb 2015

Apr 2015 Jul 2015

\$55.31

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33 May June Recap of Mylan Actions to Date April March 2015

Source of Mylan information: Mylan public filings Mylan Teva April 27: Teva reiterates commitment to Mylan proposal May 5: Teva provides additional detail on Mylan proposal and files updated investor presentation June 19: Teva announces completion of purchase of 4.61% interest in Mylan June 8: Vigodman and Peterburg write letter to Coury May 27: Teva first discloses 1.35% stake in Mylan April 29: Teva sends letter to Mylan **Board** April 24: Erez Vigodman has in-person meeting with Robert Coury April 21: Teva sends letter to Robert Coury and proposes to acquire Mylan April 29: Mylan raises offer

to acquire Perrigo

April 24: Mylan

commences

formal offer to

acquire Perrigo

April 17:

Mylan

rejects Teva

offer before a proposal

is announced

April 3:

Mylan

enters Call

Option Agreement

with the Foundation

March 10:

Market speculation of

Teva/Mylan transaction

in a Cowen research

report

April 8:

Mylan

proposes to

acquire Perrigo

April 27:

Mylan

rejects Teva

proposal in letter from

Robert Coury

June 1:

Robert Coury

writes letter to

Erez

Vigodman

June 8:

Coury

writes

response letter

to Vigodman

and Peterburg

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34 Pathway to Success Vote against proposed Perrigo transaction Obtain Board control

Because of Mylan s unprecedented board control provisions, Teva is confident that a path to Board control can be created by Dutch courts if

necessary

Obtain all applicable antitrust approvals

Teva

has already filed for U.S. HSR antitrust clearance and initiated the premerger notification process with the European Commission

Teva

has successful track record of completing transactions and working to satisfy the concerns of antitrust regulators Established Dutch methods allow for acquisition of all of Mylan

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Teva s Offer for Mylan Represents a Superior Alternative to a Mylan / Perrigo Combination
Teva s Proposal for Mylan
Mylan s Proposal for Perrigo
1.

Compared to the unaffected stock price of \$55.31 on 03/10/15, after which there was widespread speculation of a transaction be

2.

Per Mylan

offer announcement dated April 24, 2015

A clear industry leader with a larger global

manufacturing footprint and leading

positions in key product areas

Smaller scale

Weaker

strategic fit

Stronger financial profile with projected pro

forma revenue and EBITDA of almost double

that

of

Mylan

Perrigo

by

2018

Weaker financial profile and lower cash flow

generation for deleveraging

Significant \$2 billion of synergies achievable

within three years of the transaction date

Lower synergies of \$800 million achievable over

a

longer

time

horizon

of

four

years

(2)

A substantial 48% premium to Mylan s

unaffected stock price

(1)

and immediate

cash value for Mylan stockholders

Paying a premium rather than receiving one

Limited value creation for Mylan stockholders

Upside participation

No upfront liquidity for

Mylan stockholders

Teva s

proposal creates a stronger business and delivers more value to Mylan

stockholders than a Mylan

/ Perrigo

combination

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Agenda
Teva s
Business Transformation
Teva
and Mylan
Transaction Overview

Clear and Compelling Strategic Rationale
Pathway to Completion
Conclusion and Next steps
Clearly Superior Alternative to a Mylan
/ Perrigo
Combination
1
2
a
b
c
d
e

1.
Premium to Mylan unaffected price as of March 10, 2015 being the last date before there was widespread speculation of a tractive value proposition for Mylan s

stockholders

Offer price represents

a

48.3%

premium

(1)

Significant short-term value

creation and large cash

component

\$2 billion synergies drive attractive

long-term value upside

Financial strength of combined

business is a strong platform for

growth and future M&A

Compelling strategic rationale

Meaningful and real commitment

from Teva

Clear pathway to completion

Superior

alternative

to

Perrigo

for

stockholders and stakeholders

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Teva s

Offer

is

the

Superior

Outcome

for

Mylan

Stockholders

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