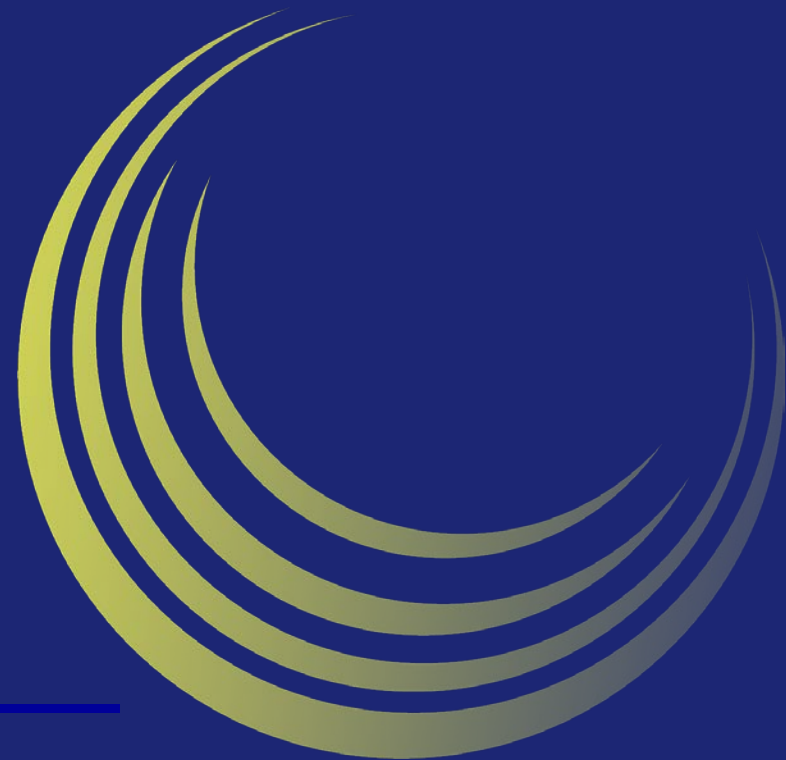


Supernus Pharmaceuticals



Investor Presentation

November 2019

Safe Harbor Statement

This presentation and other matters discussed today or answers that may be given to questions asked include forward-looking statements within the meaning of the federal securities laws. These statements, among other things, relate to Supernus' business strategy, goals and expectations concerning its product candidates, future operations, prospects, plans and objectives of management. The words "anticipate", "believe", "could", "estimate", "expect", "intend", "may", "plan", "predict", "project", "will", and similar terms and phrases are used to identify forward-looking statements in this presentation. Supernus' operations involve risks and uncertainties, many of which are outside its control, and any one of which, or a combination of which, could materially affect its results of operations and whether the forward-looking statements ultimately prove to be correct. Supernus assumes no obligation to update any forward-looking statements except as required by applicable law.

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Proven Execution in CNS & ADHD

20+ Years of CNS experience including Four Programs in ADHD

<p>2005 to Present</p>		 	<p>SPN-812 SPN-809 SPN-604 SPN-817</p>
<p>1997 to 2005</p>		 	 

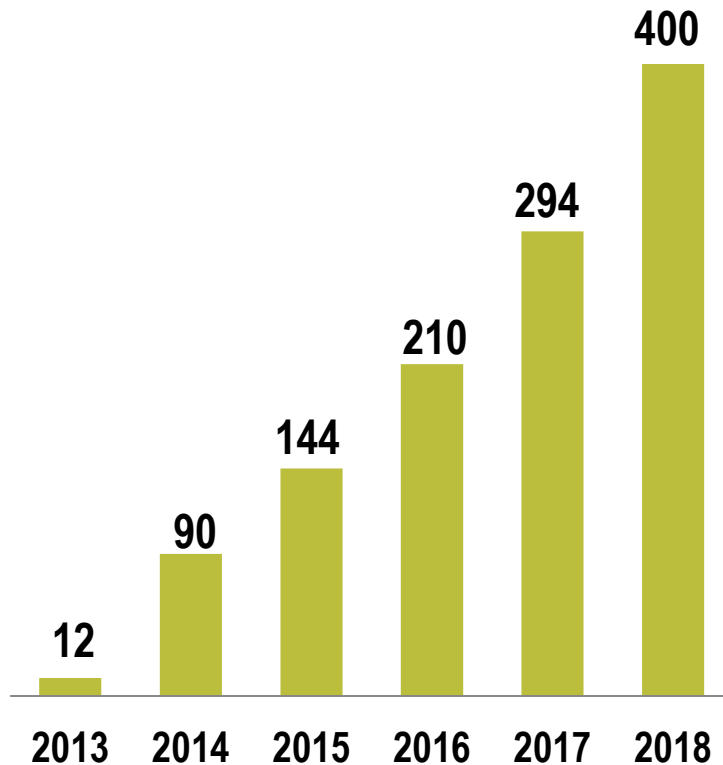
For several years, and prior to becoming independent in 2005, Supernus operated as Shire Laboratories, Inc., a division of Shire. SPN-812, SPN-810, SPN-809, SPN-604, and SPN-817 are product candidates in various stages of development. All trademarks are the property of their respective owners



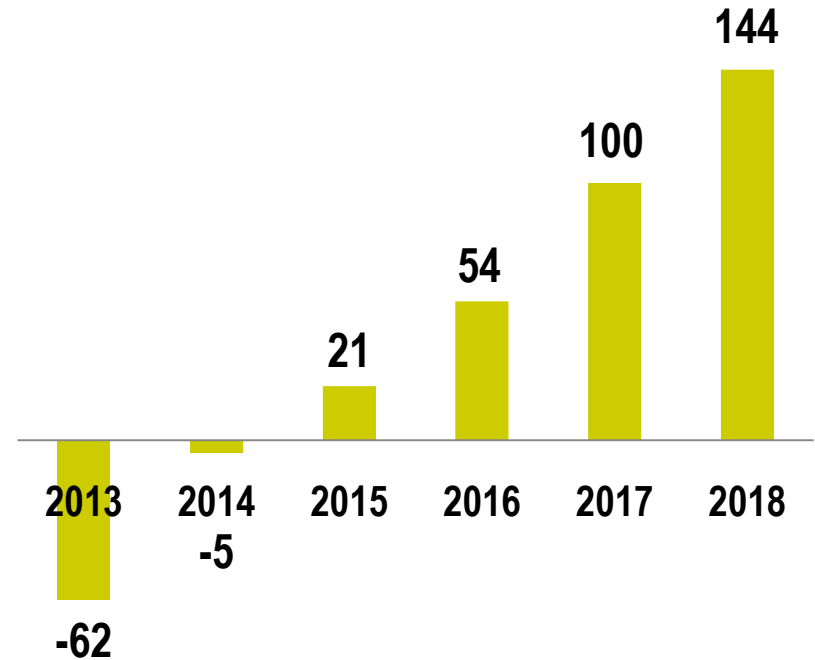
Profitable CNS Company

Strong Sales and Operating Earnings Growth

Total Net Product Sales (\$ Millions)

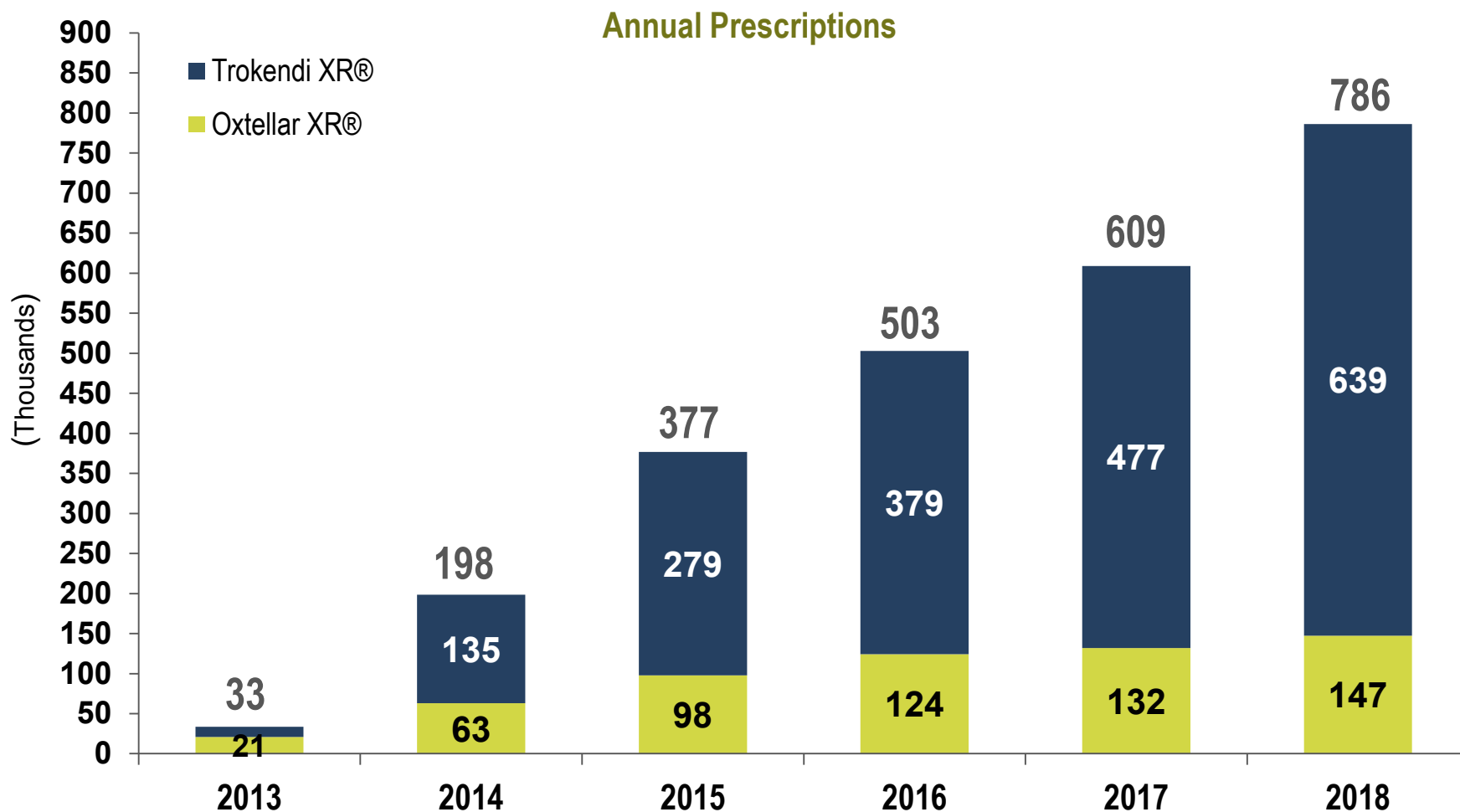


Total Operating Earnings (\$ Millions)



Trokendi XR and Oxtellar XR Prescription Growth

Combined January through October 2019 Prescription Growth of 8%



Source: IQVIA Monthly Prescriptions - Include recent restatement for 2017 and 2018 by IQVIA

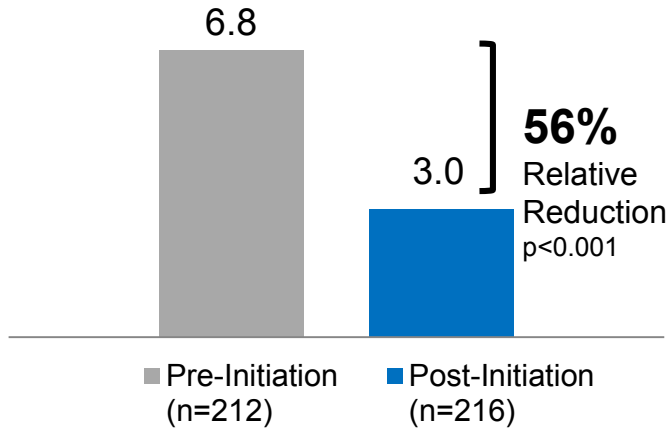
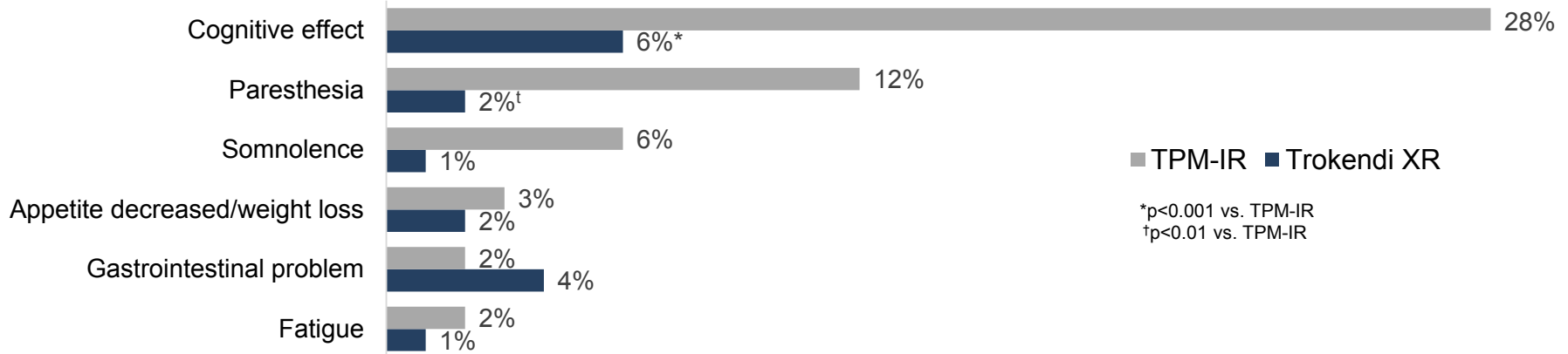
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Trokendi XR

More Favorable Clinical Outcomes Compared to TPM-IR¹

Side Effects with Trokendi XR vs. TPM-IR in Migraine Cohort (n=124)



Median Monthly Migraine Frequency
 Pre- vs. Post-Initiation of Trokendi XR

¹ O'Neal W et al. Cognitive tolerability and health outcomes with Trokendi XR (extended-release topiramate) in migraineurs. J Pain 2017; 18(4): S67. Retrospective Medical Chart Review

TPM-IR = Topiramate immediate release



Trokendi XR

Use in Clinical Practice – A Pragmatic Assessment¹

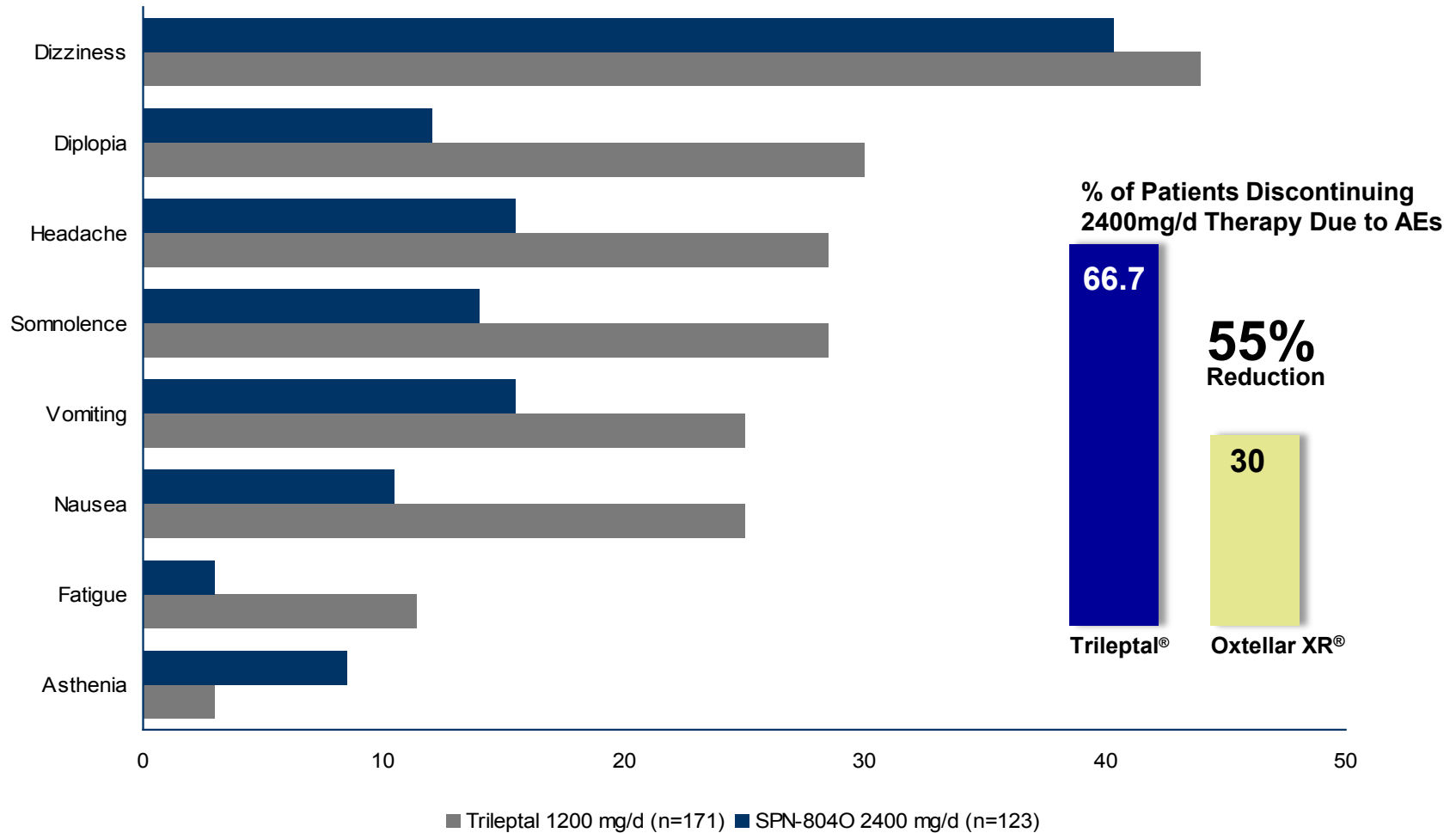
Responder Rate	% of Patients
≥ 50% Reduction	55
≥ 75% Reduction	41
100% Reduction	24

* Responder Rate: percent change from pre-index migraine frequency associated with Trokendi XR treatment (n=159)

¹ O'Neal W et al. Pragmatic assessment of Trokendi XR (extended-release topiramate) in migraine prevention. Poster presented at 59th Annual Scientific Meeting of the American Headache Society, June 2017

Oxtellar XR

Improved Adverse Event Profile at Double the Dose of Trileptal®



Based on comparison of Oxtellar XR (SPN-804O) Phase III vs. Trileptal PI (adjunctive therapy study in adults); differences in trial design exist between the two studies. Dizziness includes vertigo in Trileptal group because of change in the MedDRA system

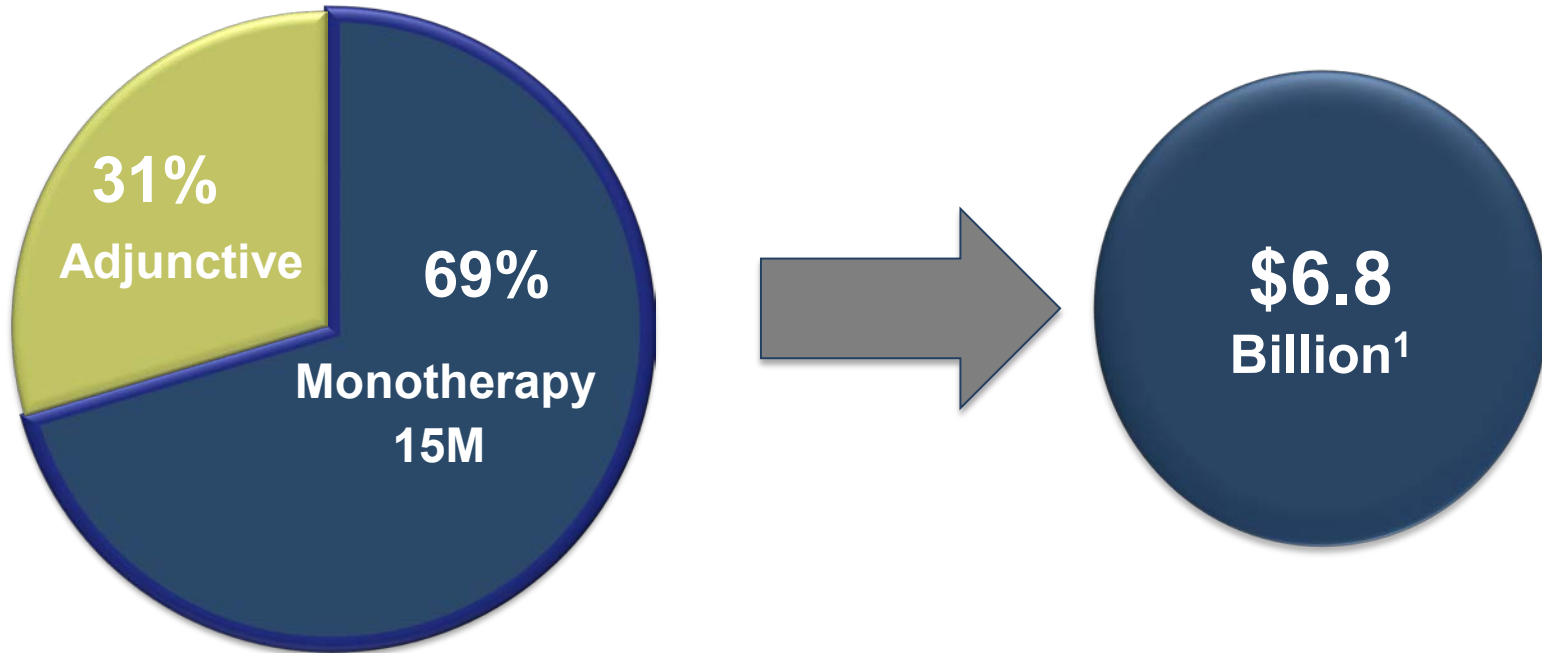
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Monotherapy Epilepsy Market Opportunity

69% of Partial Seizure Prescriptions Are For Monotherapy

Partial Seizure Prescriptions 22M Annually



Oxcarbazepine – Studied in Monotherapy with 8 Positive Clinical Trials²

IMS NDTI MAT12 months



¹ Using a branded TRx at \$450 Net

² Glauser TA. *Pharmacother.* 2001;21:904-919

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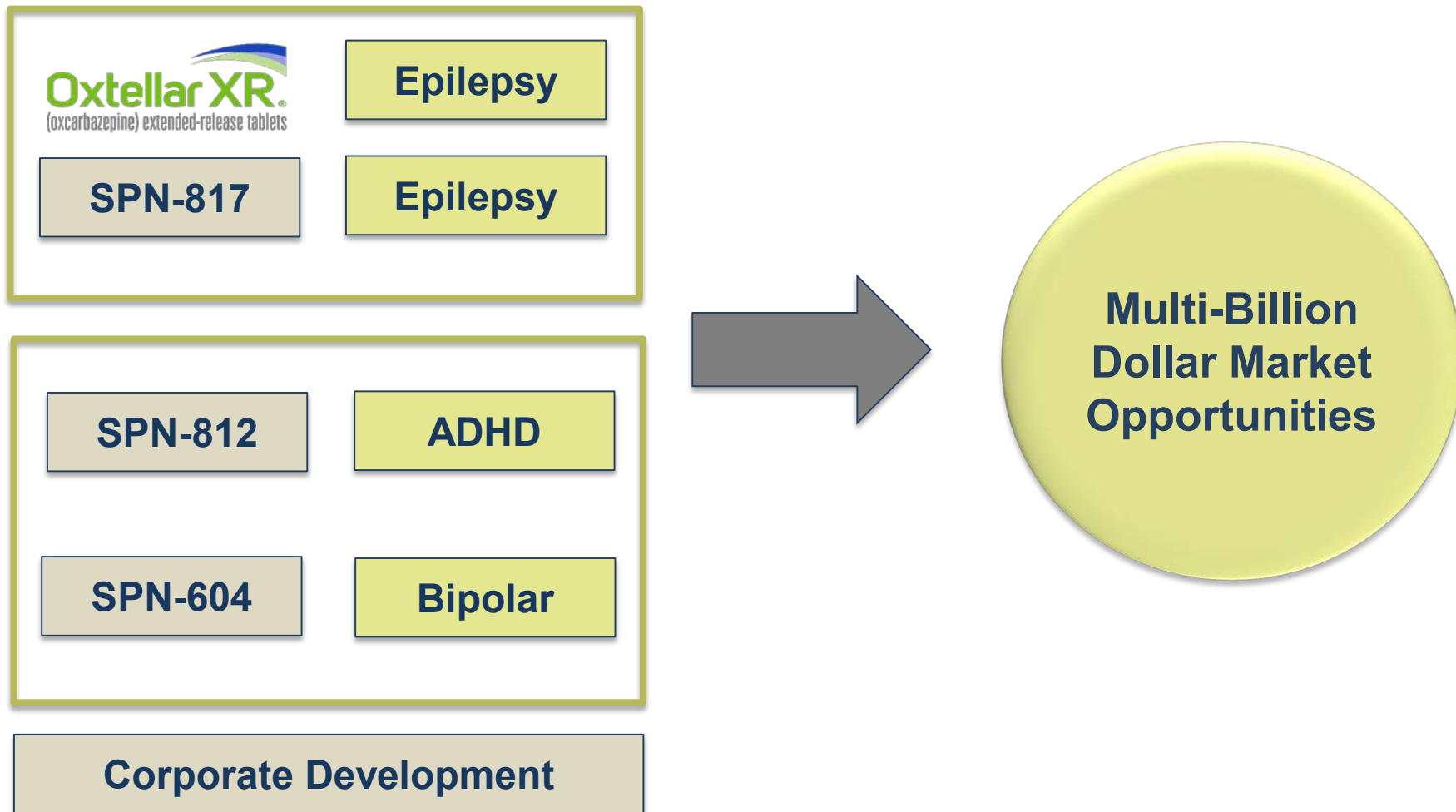
Robust Portfolio of CNS Products

Marketed	 Trokendi XR. (topiramate) extended-release capsules	Epilepsy / Migraine*		
	 Oxtellar XR. (oxcarbazepine) extended-release tablets	Epilepsy		
	Product	Indication	Development	NDA
Pipeline	SPN-812	ADHD	Phase III	Filed
	SPN-604	Bipolar	Phase III	
	SPN-809	Depression	IND/Phase II Ready	
	SPN-817	Severe Epilepsy	Phase I	

*Prophylaxis of migraine in adolescents and adults

Future Growth Drivers

Several Significant Market Opportunities



SPN-812

Novel Non-Stimulant ADHD Product Candidate

- Viloxazine hydrochloride
 - Serotonin norepinephrine modulating agent (SNMA)
 - New Chemical Entity (NCE)
 - Previously marketed outside the US as an antidepressant
- Building strong IP with expirations from 2029-2033
- NDA submitted to FDA in November 2019
- Phase III clinical data point to a well-differentiated ADHD product
 - 100mg, 200mg and 400mg in pediatric patients
 - Unique mechanism of action
 - Consistent & reliable efficacy with robust statistical significance
 - Efficacy on both Hyperactivity/Impulsivity and Inattention
 - Fast onset of action
 - Well tolerated

SPN-812

Phase III Studies

	P301 N = 477	P303 N = 313	P302 N = 310	P304 N = 297
ADHD Patients	6-11 years	6-11 years	12-17 years	12-17 years
Daily Doses	100mg 200mg	200mg 400mg	200mg 400mg	400mg 600mg
Status	Completed	Completed	Completed	Completed

Randomized, double-blind, placebo-controlled, multicenter, parallel group, monotherapy for ADHD
 Primary Endpoint - Change from baseline on ADHD-RS-5 scale compared to placebo



SPN-812 Phase III Data: Primary Endpoint

P301 (Children)	Statistics	Placebo (N=155)	100 mg (N=147)	200 mg (N=158)
Week 6 (EOS)	LS Mean	-10.9	-16.6	-17.7
	p-value		0.0004	<.0001
P302 (Adolescent)	Statistics	Placebo (N=104)	200 mg (N=94)	400 mg (N=103)
Week 6 (EOS)	LS Mean	-11.4	-16.0	-16.5
	p-value		0.0232	0.0091
P303 (Children)	Statistics	Placebo (N=97)	200 mg (N=107)	400 mg (N=97)
Week 8 (EOS)	LS Mean	-11.7	-17.6	-17.5
	p-value		0.0038	0.0063
P304 (Adolescent)	Statistics	Placebo (N=97)	400 mg (N=99)	600 mg (N=97)
Week 7 (EOS)	LS Mean	-13.2	-18.3	-16.7
	p-value		0.0082	0.0712

Primary Analysis of ADHD-RS-5 based on Mixed Model for Repeated Measure (MMRM) Intent to Treat (ITT Population)
EOS = End of Study



SPN-812 Phase III Data

Significant Reduction in Hyperactivity and Inattention

Analysis in ADHD-RS-5 Inattention and Hyperactivity/Impulsivity Subscales

P301 Week 6 (EOS)	Statistics	100 mg (N=147)	200 mg (N=158)
Hyperactivity/Impulsivity	p-value	0.0026	<.0001
Inattention	p-value	0.0006	<.0001
P302 Week 6 (EOS)	Statistics	200 mg (N=94)	400 mg (N=103)
Hyperactivity/Impulsivity	p-value	0.0069	0.0005
Inattention	p-value	0.0424	0.0390
P303 Week 8 (EOS)	Statistics	200 mg (N=107)	400 mg (N=97)
Hyperactivity/Impulsivity	p-value	0.0020	0.0039
Inattention	p-value	0.0087	0.0248
P304 Week 7 (EOS)	Statistics	400 mg (N=99)	600 mg (N=97)
Hyperactivity/Impulsivity	p-value	0.0484	0.2084
Inattention	p-value	0.0042	0.1392

EOS = End of Study



SPN-812 Phase III Data: Fast Onset of Action

Efficacy Starting in Week 1 - ADHD-RS-5 Total Score

Pooled Data – P301, P302, P303, P304				
Visit	Statistics	Placebo (N=452)	200 mg (N=359)	400 mg (N=299)
Baseline	Mean	41.8	42.9	41.8
Week 1	p-value		0.0003	0.0016
Week 2	p-value		<.0001	<.0001
Week 3	p-value		<.0001	<.0001
Week 4	p-value		<.0001	<.0001
Week 5	p-value		<.0001	<.0001
Week 6	LS Mean	-11.7	-17.1	-17.7
	p-value		<.0001	<.0001

P301	
Placebo (N=155)	100 mg (N=147)
43.6	45.0
	0.0004
	<.0001
	<.0001
	<.0001
	0.0006
-10.9	-16.6
	0.0004

- Common endpoint visit for all four studies is Week 6
- Pooled Data exclude 100 mg and 600 mg that were tested in one study only
- Primary Analysis of ADHD-RS-5 in Intent to Treat Population



SPN-812 Phase III Data: Fast Onset of Action

Efficacy Starting in Week 1 - Inattention Subscale

Pooled Data – P301, P302, P303, P304				
Visit	Statistics	Placebo (N=452)	200 mg (N=359)	400 mg (N=299)
Baseline	Mean	22.4	22.6	22.3
Week 1	p-value		0.0086	0.0162
Week 2	p-value		0.0001	<.0001
Week 3	p-value		<.0001	<.0001
Week 4	p-value		<.0001	<.0001
Week 5	p-value		<.0001	<.0001
Week 6	LS Mean	-6.4	-8.9	-9.2
	p-value		<.0001	<.0001

P301	
Placebo (N=155)	100 mg (N=147)
22.5	22.8
	0.0016
	0.0016
	0.0002
	<0.0001
	0.0018
-5.6	-8.6
	0.0006

- Common endpoint visit for all four studies is Week 6
- Pooled Data exclude 100 mg and 600 mg that were tested in one study only
- Primary Analysis of ADHD-RS-5 in Intent to Treat Population



SPN-812 Phase III Data: Fast Onset of Action

Efficacy Starting in Week 1 - Hyperactivity/Impulsivity Subscale

Pooled Data – P301, P302, P303, P304				
Visit	Statistics	Placebo (N=452)	200 mg (N=359)	400 mg (N=299)
Baseline	Mean	19.4	20.3	19.5
Week 1	p-value		<.0001	0.0010
Week 2	p-value		<.0001	<.0001
Week 3	p-value		<.0001	<.0001
Week 4	p-value		<.0001	<.0001
Week 5	p-value		<.0001	<.0001
Week 6	LS Mean	-5.4	-8.2	-8.5
	p-value		<.0001	<.0001

P301	
Placebo (N=155)	100 mg (N=147)
21.1	22.2
	0.0023
	<0.0001
	<0.0001
	0.0004
	0.0010
-5.3	-8.0
	0.0014

- Common endpoint visit for all four studies is Week 6
- Pooled Data exclude 100 mg and 600 mg that were tested in one study only
- Primary Analysis of ADHD-RS-5 in Intent to Treat Population

SPN-812 Phase III Data: Secondary Endpoint

Analysis of Observed Global Improvement Score (CGI-I) at EOS

P301	Statistics	Placebo (N=155)	100 mg (N=147)	200 mg (N=158)
Week 6 (EOS)	LS Mean	3.1	2.7	2.6
	p-value		0.0020	<.0001
P302	Statistics	Placebo (N=104)	200 mg (N=94)	400 mg (N=103)
Week 6 (EOS)	LS Mean	3.0	2.5	2.4
	p-value		0.0042	0.0003
P303	Statistics	Placebo (N=97)	200 mg (N=107)	400 mg (N=97)
Week 8 (EOS)	LS Mean	3.1	2.6	2.6
	p-value		0.0028	0.0099
P304	Statistics	Placebo (N=96)	400 mg (N=99)	600 mg (N=97)
Week 7 (EOS)	LS Mean	2.9	2.4	2.6
	p-value		0.0051	0.0995

EOS = End of Study

SPN-812

Summary of Treatment Related Adverse Events

Number (%) of Patients - Treatment Related AEs with $\geq 5\%$ Incidence
All Four Phase III Trials

	Placebo (N=463)	SPN-812 (N=925)
Somnolence	14 (3.0)	115 (12.4)
Decreased appetite	2 (0.4)	61 (6.6)
Headache	14 (3.0)	57 (6.2)
Fatigue	10 (2.2)	56 (6.1)
Discontinuation due to AEs	6 (1.3)	32 (3.5)

AEs = Adverse Events

SPN-812

Significant Market Opportunity

	Percent	Estimated Prescriptions in Peak Year
ADHD Market Prescriptions		89 - 100 Million
	Peak Market Share	SPN-812 Potential Prescriptions
SPN-812 Peak Demand	5 - 10%	4.5 - 10.0 Million

Source: IMS NPA, Company Research and Estimates – Assumes peak at 3-7 years post launch
Figures in the table above represent management's estimates that are subject to several factors that are beyond our control and actual results may be significantly different from our estimates

SPN-604

Novel Product Candidate for Bipolar

50% Use of Oxcarbazepine
in Psychiatry

1st Expected to be Only
Oxcarbazepine Product
Approved to Treat Bipolar

2019 Phase 3 Program
Initiated



Market Opportunity
+53 Million Prescriptions

Class of Drugs	% of Prescriptions
Antiepileptics	34
Antipsychotics	29
SSRI's	15
SNRI's	6
Antimania	6
Other Antidepressants	6
Benzodiazepines	4
Total	100

Source: IQVIA 2016

SSRI = Selective serotonin reuptake inhibitor
SNRI = Serotonin & norepinephrine reuptake inhibitor

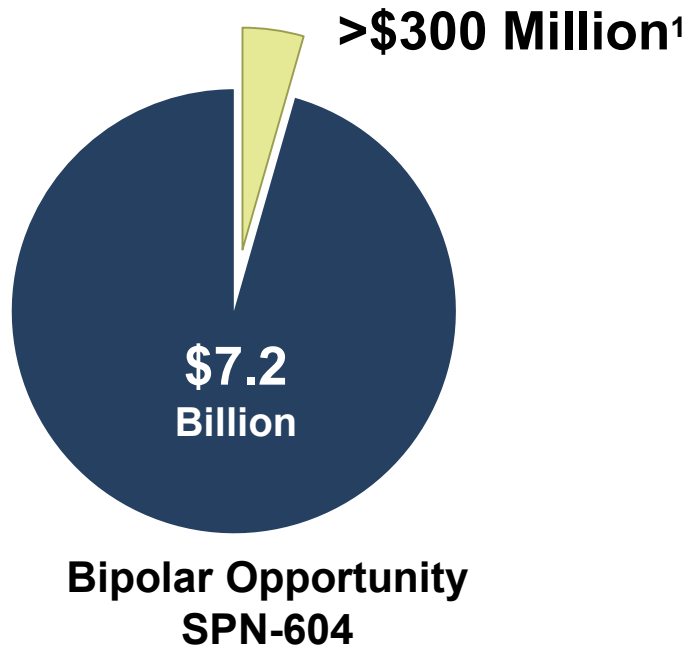
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SPN-604

Target Market Opportunity in Psychiatry of \$7.2 Billion

Potential Peak Sales - SPN-604 >\$300 Million



1- Anti-epileptic drugs represent 34% of 53 million prescriptions for bipolar (IQVIA). Average net price per prescription of \$400. Peak share of ~5%. Above figures represent management's estimates that are subject to several factors that are beyond our control and actual results may be significantly different from our estimates

Financial Summary and Guidance

3Q 2019 Financial Results

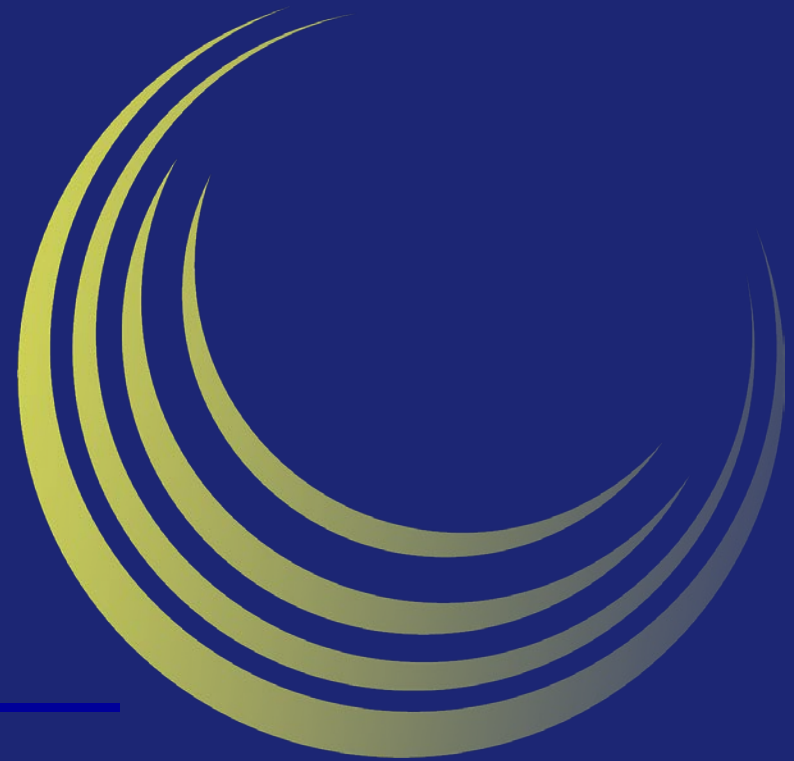
- Net sales of \$100.0 million
- Operating earnings of \$39.7 million
- Cash, cash equivalents, & investments at \$893 million as of Sept 30, 2019

Full Year 2019 Financial Guidance¹

- Net sales: \$390 million - \$395 million
- Operating earnings: \$150 million - \$155 million
 - R&D expenses: ~\$70 million

¹ Guidance as provided on November 5, 2019, and which has not been updated.

Positioned For Continued Growth in CNS



Strong Presence in Neurology with Existing Products

Oxtellar XR[®] and Trokendi XR[®]

Innovative Late Stage Portfolio in Psychiatry

SPN-812

Well Differentiated Novel Non-Stimulant

SPN-604

Novel Product for Bipolar Disorder