# Budget Impact of Somatostatin Analogs (SSAs) as Treatment for Metastatic Gastroenteropancreatic Neuroendocrine Tumors (mGEP-NETs) in US Hospitals Jesse D. Ortendahl<sup>1</sup>; Sonia J. Pulgar<sup>2</sup>; David Cox<sup>2</sup>; Tanya G.K. Bentley<sup>1</sup>; Alexandria T. Phan<sup>3</sup> <sup>1</sup>Partnership for Health Analytic Research; <sup>2</sup>Ipsen Biopharmaceuticals; <sup>3</sup>Houston Methodist Hospital

### BACKGROUND

- Gastroenteropancreatic neuroendocrine tumors (GEP-NETs) are rare neoplasms that originate in the secretory cells of the neuroendocrine system. Many of these tumors produce peptides and neuroamines, causing characteristic hormonal syndromes such as carcinoid syndrome.<sup>1,2</sup>
- Diagnosed NETs incidence is increasing, with US prevalence likely exceeding 100,000.<sup>3,4</sup>
- Initial systemic therapy for NETs often consists of treatment with a somatostatin analog (SSA) such as lanreotide depot (Somatuline) and octreotide LAR (Sandostatin).

### **OBJECTIVE**

- In the United States, only lanreotide depot is indicated in patients with GEP-NETs to improve progression free survival; octreotide LAR is approved for the treatment of severe diarrhea/ flushing episodes associated with metastatic carcinoid tumors. Both products are included in guidelines and are used frequently in practice.
- Trials have shown improved progression-free survival with both products when compared with placebo.<sup>5,6</sup>
- Real world costs of SSAs can be influenced by how they are used in clinical practice. Budget impact analysis (BIA) can be useful because it considers how changes in the mix of drugs and factors such as dosing, efficiency, and injection success can impact the trajectory of healthcare spending within a specific healthcare setting. This BIA is based on cost and dosing within the therapeutic class and does not evaluate efficacy or safety outcomes.<sup>7</sup>

This model-based budget impact analysis estimates the financial impact on US hospitals of adopting and shifting the mix of somatostatin analogs (SSAs) utilized to treat GEP-NETs.

### **METHODS**

#### Model Overview

- Structure: Deterministic cohort model
- **Population:** Patients with metastatic GEP-NETs
- **Perspective:** US hospital
- Time horizon: 1 year

#### Model Structure

- Patients entered the model and received either lanreotide depot or octreotide LAR.
- The model assessed two scenarios:
  - Current Utilization, in which the market share reflects current treatment patterns;

Table 4: Market Share by Product <sup>a</sup>		
Droduct	Current	Comparison
FIUUUCI	Current Utilization	Scenario
Lanreotide Depot	5%	30%
Ostrastida I A D	050/	700/

Model inputs: Patients eligible for SSA treatment, product acquisition costs, preparation and mixing costs, product utilization Outcome measures: Annual costs, costs per treated patient

Parameter	Value	Source
GEP-NET Patients (N)	500	Assumption
Metastatic/Inoperable	80.0%	8
Treated with an SSA	78.2%	9
<b>Final Model Patient Population (N)</b>	313	

GEP-NET, gastroenteropancreatic neuroendocrine tumor; SSA, somatostatin analogue.

Table 2. Product Acquisition Costs <sup>a</sup>	
Product	Acquisition Cost (per syringe) <sup>10</sup>
Lanreotide Depot	
60mg	\$3,328
90mg	\$4,434
120mg	\$5,494
Octreotide LAR	
10mg	\$2,380
20mg	\$3,118
30mg	\$4,670
<sup>a</sup> Prices reported as wholesale acquisition costs	s (WAC).

## RESULTS

#### **Base Case Results**

In the base case, lanreotide depot reduced per-patient costs compared with octreotide LAR (\$71,442 vs. \$75,508).

- Comparator Scenario, which reflects a hypothetical shift in utilization from octreotide LAR to lanreotide depot.
- The model population was based on a hypothetical hospital with 500 GEP-NET patients, assuming 80% had metastatic disease or were inoperable, and 78% were treated with an SSA (Table 1).
- Patients were assumed to be treated for the entirety of the 1year period with 100% medication adherence.
- Dosing patterns and injection frequency of each product remain constant for the two scenarios.
- Patients incurred costs associated with product acquisition and administration (Tables 2-3).

Table 3. Product Pre	eparation and Administrat	tion Costs	
Product	Mixing/ Preparation Time (seconds) <sup>11</sup>	Wage Rate <sup>12</sup>	Cost (per syring
Lanreotide Depot	66	<b>¢01 17</b>	\$1.51
Octreotide LAR	329	\$82.27	\$7.52

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L	<sup>a</sup> Current utilization scenario defined as current marke	t share based on n	narket research data.
	Comparison scenario defined as a hypothetical shift in	market share.	

Table 5: Distribution	n of Dosing by Pro	duct <sup>a</sup>	
Product	1x per 4 weeks	1x per 3 weeks	1x per 2 weeks
Lanreotide Depot			
60mg	0%	0%	0%
90mg	0%	0%	0%
120mg	100%	0%	0%
Octreotide LAR <sup>13</sup>			
10mg	0%	0%	0%
20mg	0%	0%	0%
30mg	64%	3%	7%
40mg	14%	6%	0%

0% 2% 4% 60mg <sup>a</sup> Default estimates for lanreotide depot based on the labeled dosing regimen of 120 mg every four weeks. For octreotide LAR, proportion receiving dosing above indication based on a real-world study. All others assumed to get highest indicated dose.

#### Analyses

- **Base case:** Total hospital and per patient costs were estimated for the current utilization and the comparison scenario.
- One-way sensitivity analyses: All model parameters were independently varied by +/- 20%.

• For a hypothetical hospital with 500 GEP-NET patients, the annual cost of shifting from 5% to 30% lanreotide depot use resulted in cost savings of \$317,977 (\$23,555,246 vs. \$23,237,269).

• In one-way sensitivity analyses, results were shown to be driven by product acquisition costs and proportion of octreotide LAR patients receiving above indicated dosing.

	Annual Hospital/I	Annual Hospital/Institution Costs		Cost per Treated Patient		
	Current Utilization	Comparison Scenario	Product Acquisition	Preparation and Mixing	Total	
anreotide depot	\$1,117,347	\$6,704,080	\$71,422	\$20	\$71,442	
ctreotide LAR	\$22,437,899	\$16,533,189	\$75,400	\$108	\$75 <i>,</i> 508	
otal	\$23,555,246	\$23,237,269	-	-	-	
ifference <sup>c</sup>	_	-\$317,977	_	-	-	

<sup>c</sup> Difference reflects the change in total costs between the baseline and comparator year. A negative number denotes a cost savings in comparator year.



### REFERENCES

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# Conclusions

Results were most sensitive to the acquisition costs for each product and the proportion of octreotide patients receiving above indicated dosing.

- Results from this analysis suggest that factors beyond drug acquisition cost can influence the overall hospital budget impact with SSA treatment for GEP-NETs. • Incorporating factors such as higher than indicated dosing of octreotide LAR observed in a real-world study and differential preparation and mixing costs, we found that increasing use of lanreotide depot at its indicated dosing results in cost-savings for the hospital.
- These results are based on the current treatment patterns of SSAs in the US; these results may change as more real world data becomes available, especially for lanreotide depot, which became available more recently in the US for management of NETs.

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