#### **Public observer slides**

# Everolimus, lutetium-177 DOTATATE and sunitinib for treating unresectable or metastatic neuroendocrine tumours with disease progression – MTA

1<sup>st</sup> Appraisal Committee meeting

Background and Clinical Effectiveness

Committee D

Lead team: Dr Ian Davidson, Gillian Ells and Pam Rees

ERG: Peninsula Technology Assessment Group

NICE technical team: Stuart Wood and Nwamaka Umeweni

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### Key clinical issues

- How are NETs treated in clinical practice?
- Have the appropriate comparisons been made for each tumour location?
- Can the results from pivotal trials be generalised to current clinical practice?
- What conclusions can be drawn from the network meta-analyses
- Are everolimus and sunitinib clinically equivalent?

# Neuroendocrine tumours (NETs)

- Heterogeneous group of rare tumours that develop from the gastrointestinal tissue, pancreas, lung and thyroid
- Approximately 45-65% of NETs occur in the gastrointestinal tissue, approximately 3-7% in the pancreas and 10% in the lungs
- Can be 'functional' or 'non-functional'
- Grade of the tumour gives an idea of how quickly it will develop low (grade 1) well differentiated moderate (grade 2) high grade tumours (grade 3) poorly differentiated
- Ki-67 proliferative index (Ki-67 index) is also used as a prognostic measure for grading parameters for NETs

Grade 1 is equivalent to a Ki67 index of up to 3%

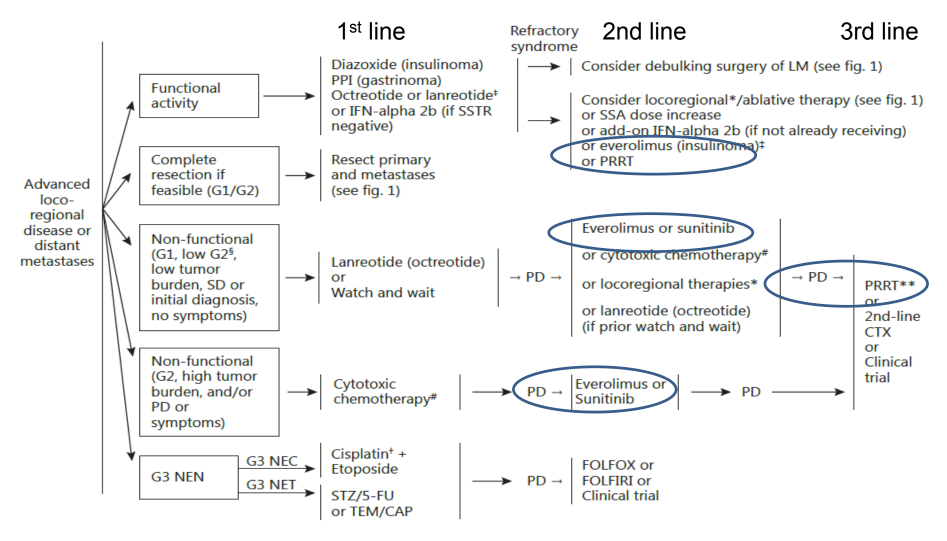
Grade 2 is equivalent to a Ki67 index between 3-20%

Grade 3 is equivalent to a Ki67 index beyond 20%

# Neuroendocrine tumours (Management)

- No NICE guidance on neuroendocrine tumours
- Surgery is the only curative treatment
- Options for treating progressed neuroendocrine tumours include:
  - Somatostatin analogues (for symptomatic control e.g. octreotide, lanreotide)
  - Chemotherapy regimens (using combinations of streptozocin, 5fluorouracil, doxorubicin, temozolomide and capecitabine)
  - Radionuclides (e.g. lutetium-177 previously on the CDF)
  - Everolimus (previously on the CDF)
  - Sunitinib (currently on the CDF)
- Limited data for lung NETs
  - ENETS guidelines recommends everolimus for progressive lung NETs

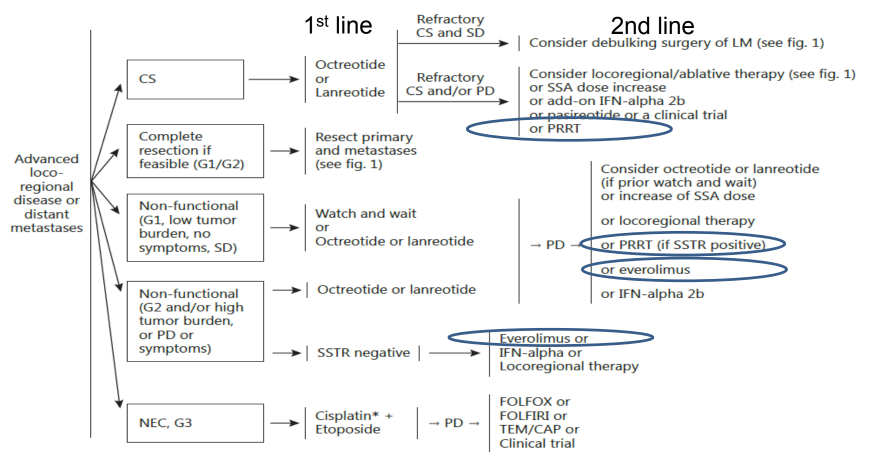
# Treatment pathway: Pancreatic NETs



Source: Novartis submission, figure 3.1, page 29

Original source: Pavel et al 2016, ENETs-recommended treatment algorithm

# Treatment pathway: GI NETs



Abbreviations: 5-FU: 5-fluorouracil, CAP: capecitabine, CS: carcinoid syndrome, CTX: chemotherapy, FOLFIRI: folinic acid, 5-FU, irinotecan, FOLFOX: folinic acid, 5-FU, oxaliplatin, IFN: interferon, LM: liver metastases, NEN: neuroendocrine neoplasm, PD: progressive disease, PRRT: peptide receptor radionuclide therapy, SD: stable disease, SSA: somatostatin analogue, SSTR: somatostatin receptor, STZ: streptozotocin, TEM: temozolomide.

Source: Pavel et al. 2016

Source: Novartis submission, figure 3.1, page 29

Original source: Pavel et al 2016, ENETs-recommended treatment algorithm

# Patient perspectives

#### **NET Patient Foundation**

- Challenging tumours to diagnose and treat
- Around 3,000 new diagnoses each year in UK, but many remain undiagnosed
- Historically, treatments often improved symptoms but not always overall survival
- New treatments have improved progression-free survival, but also increased toxicity
- High unmet need in patients with lung NETs, and patients with GI NETs who have progressed following current therapy
- No NICE guidance
- Patients in England have seen their options increasingly restricted over the past two years
- PRRT and the Cancer Drug Fund

# Patient perspectives (2)

#### **NET Patient Foundation**

- 'No clear pathway' of care for patient with NETs
- Lack of clarity and certainty impact on HRQoL & wellbeing
- Results from First Global NET survey:
  - 60% patients reported NETS negatively impacted emotional health
  - 52% experience significant stress & anxiety levels
  - 39% feel confused about the management of their disease
  - Of the 22% who were not working/unemployed due to medical disability,
     82% had stopped working as a result of their NET
- Patients experience of Lu177 DOTATATE has been positive with significant improvement to length of life and quality of life

# Clinical perspectives (1)

#### British Institute of Radiology and British Nuclear Medicine Society

- Majority of well differentiated NETS express somatostatin receptors on their surface which can be targeted by somatostatin receptor based radionuclide therapy
- Lu-177 DOTATATE is an effective treatment and place in treatment algorithms is recommended by several international guidelines including ENETS Consensus Guidelines (2016)
- Lu-177 DOTATATE is promoted as second-line therapy for disease progression after first-line therapy with SSA's
- The guidelines also recommend its use as third-line therapy after everolimus in non-midgut NETs
- In patients with progressive disease Lu-177 DOTATATE stabilises disease and prolongs survival and side effects are uncommon
- QoL analysis in 39 consecutive patients at the Royal Free London NHS Foundation Trust demonstrated a significant improvement in QOL in patients treated with Lu-177 DOTATATE in NETs
- Lu-177 DOTATATE is a safe and efficacious treatment for metastatic NETs
- Number of centres in the UK already providing Lu177 DOTATATE
- No further resources would be required for provision of Lu177 DOTATATE

# Clinical perspectives (2)

#### Royal College of Physicians

- Management of NETs requires a multi-disciplinary treatment approach
- Limiting to patients with advanced disease and well-differentiated is appropriate
- Disease progression, treatment choice depends on site of tumour
- In P-NETs, everolimus and sunitinib are a clinically effective treatment option giving patients extra lines of therapy
  - Treatment continues until progression
- In intestinal NETs, treatment options beyond SSA's are limited
- 177-Lu DOTATATE allows the use of targeted radiotherapy, likely to be effective in all NETs
- No recognised optimal sequence of therapies
- Targeted therapies have the same level of activity regardless of prior chemotherapy use
- Clinicians are familiar with everolimus and sunitinib and management of toxicity
  - Adverse events are manageable
- NICE positive guidance would allow the UK to remain as one of the leading countries in NET patient-centred care

### **DETAILS OF THE TECHNOLOGIES**

	Lutetium-177 DOTATATE (Lutathera, AAA)	Everolimus (Afinitor, Novartis)	Sunitinib (Sutent, Pfizer)
MA		<ul> <li>unresectable or metastatic, well- or moderately-differentiated neuroendocrine tumours of pancreatic origin in adults with progressive disease</li> <li>unresectable or metastatic, well-differentiated (grade 1 or grade 2) non-functional neuroendocrine tumours of gastrointestinal or lung origin in adults with progressive disease</li> </ul>	unresectable or metastatic, well-differentiated pancreatic neuroendocrine tumours with disease progression in adults
Admin.	Intravenous Infusion (IV)	Oral	Oral
Costs	<ul> <li>A single cycle comprising four administrations of 7.4 GBq. The recommended interval between two infusions is eight weeks (± 1 week).</li> </ul>	<ul> <li>The list price for everolimus is £2,673.00 for 30 x 10 mg everolimus tablets</li> <li>A confidential PAS is available and details are presented in a confidential appendix</li> </ul>	<ul> <li>Pack of 28, 12.5 mg capsules £784.70.</li> <li>Pack of 29, 25 mg capsules £1,569.40.</li> <li>Pack of 28, 50 mg capsules £3,138.80.</li> </ul>

# **DECISION PROBLEM**

health-related quality of life

Final s	cope issued by NICE	AG comments
Pop	People with progressed unresectable or metastatic neuroendocrine tumours (according to the specific locations covered by the existing and anticipated marketing authorisations of the interventions)	The AG population is consistent with the NICE scope
Int	<ul> <li>Everolimus (GI, Pancreatic or Lung NETS)</li> <li>Lutetium-177 DOTATATE (GI or Pancreatic NETs)</li> <li>Sunitinib (Pancreatic NETs)</li> </ul>	The AG included all of these interventions
Comp	<ul> <li>the technologies listed above will be compared with each other where appropriate</li> <li>interferon alpha</li> <li>chemotherapy regimens</li> <li>best supportive care</li> </ul>	The AG consulted with clinicians and were told that interferon alpha was not commonly used within UK clinical practice. Therefore, it was not included
Out	<ul> <li>overall survival</li> <li>progression-free survival</li> <li>response rates</li> <li>symptom control</li> <li>adverse effects of treatment</li> </ul>	The AG considered and included all of these outcome measures

#### Pancreatic NETs: Clinical Trials

Trial	RADIANT-3 – everolimus plus BSC Vs. placebo plus BSC	A6181111 – sunitinib plus BSC Vs. placebo plus BSC	
Design	Double-blind, randomised, placebo-controlled phase III		
Population	<ul> <li>Patients with advanced, progressive, low- or intermediate- grade P-NETs</li> </ul>	<ul> <li>Patients with progressive well-differentiated P-NETs</li> </ul>	
Outcomes	<ul> <li>Primary endpoint - PFS (locally assessed according to RECIST)</li> <li>Secondary endpoints - OS, DoR, ORR and safety</li> </ul>	<ul> <li>Primary endpoint – PFS</li> <li>Secondary endpoints - OS, ORR, TTR, DoR, EORTC QLQ-C30 (HRQoL)</li> </ul>	
Other	<ul> <li>Concurrent SSA use allowed (37.7 % and 39.9% in the everolimus and placebo arms respectively)</li> <li>Crossover from the placebo arm to the treatment arm was 73%</li> </ul>	<ul> <li>SSA use permitted both before and during the trial</li> <li>Cross-over allowed (at disease progression) in one of two separate, open-label extension studies</li> <li>69% placebo patients crossed over to sunitinib</li> </ul>	

# **RADIANT-3 Results**

Novartis submission, tables $4.3 - 4.5$ (pages $37 - 44$ )					
Outcomes Local assessment Adjudicated central r					
Progression-free survival (PFS)					

Placebo + BSC

(n=203)

4.6

(3.1 - 5.4)

Overall survival (OS) with adjustment for cross-over (Final OS analysis, March 2014, open

37.68

4 (2.0)

**103** (50.7)

**85** (41.9)

Everolimus +

**BSC** 

(n=207)

11.4

the local review

to RECIST

Compared with placebo,

Everolimus +

**BSC** 

(n=207)

11.0

(8.4 - 13.9)

44.02

**10** (4.8)

**151** (72.9)

**29** (14.0)

**0.35** (0.27–0.45)

**0.60** (0.09–3.95)

PFS, median,

HR (95% CI)

label phase)

OS. median.

HR (95% CI)

Partial response

Stable disease

Progressed

disease

**Tumour response rates (n%)** 

months

months

W

Placebo + BSC

(n=203)

5.4

0.34 (0.26 - 0.44)

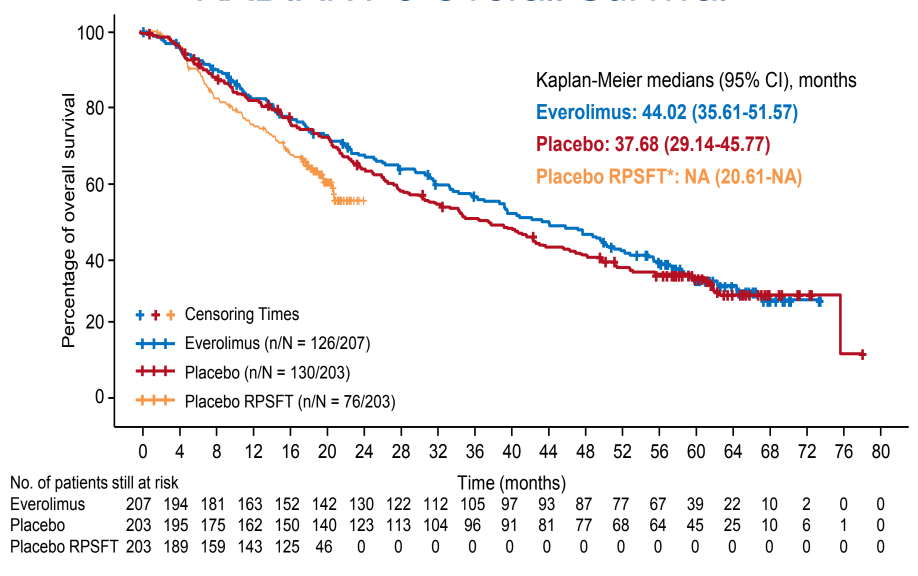
Results from the central reviews

were similar to those reported for

everolimus was associated with a

superior response profile according

### **RADIANT-3 Overall Survival**



Source: Novartis submission, figure 4.7, page 45

### RADIANT-3 subgroup analyses

PFS subgroup analysis				
Covariate	Covariate Subgroup		HR (95% CI)	
	Well differentiated	341	<b>0.41</b> (0.31, 0.53) P<0.001	
Tumour grade:	Moderately differentiated	65	<b>0.21</b> (0.11, 0.42) P<0.001	
Previous	Yes	189	<b>0.34</b> (0.24,0.49) P<0.001	
chemotherapy	No	221	<b>0.41</b> (0.29,0.58) P<0.001	
Previous long-acting	Yes	203	<b>0.40</b> (0.28,0.57) P<0.001	
SSA use	No	207	<b>0.36</b> (0.25,0.51) P<0.001	

OS subgroup analysis				
Covariate	Subgroup	N	HR (95% CI)	
Previous	Yes	189		
chemotherapy	No	221	<b>0.78</b> (0.61, 1.01) P=0.056	
Previous long-acting	Yes	203		
SSA use	No	207	<b>1.15</b> (0.89, 1.49) P=0.288	

Sources: Assessment report, tables 29 – 30, page 96

### A6181111 Results (1)

Pfizer submission, section 4.7, pages 42 - 50

Outcomes	Investigator	assessment	Independent review	
	Sunitinib (n=86)	Placebo (n=85)	Sunitinib (n=86)	Placebo (n=85)
Progression-free	survival			
PFS, median, months	<b>11.4</b> (7.4 – 19.8)	<b>5.5</b> (3.6 – 7.4)	<b>12.6</b> (11.1 - 20.6)	<b>5.8</b> (3.8 - 7.2)
HR (95% CI)	<b>0.418</b> (CI: 0	.263, 0.662)	<b>0.315</b> (0.1	81, 0.546)
Overall survival				
OS unadjusted for cross over,	<b>38.6</b> (25.6 – 56.4)	<b>29.1</b> (16.4 – 36.8)		_
median, months HR (95% CI)	<b>0.73</b> (0.5	50 – 1.06)		
Adjustment for crossover, median, months – RPSFT (placebo)	-	<b>13.2</b> (11.3 – 16.5) <b>HR 0.34</b> (0.14 – 1.28)	-	_
Censoring at crossover – IPCW (placebo)	-	<b>16.3</b> (12.5 – 24.3) <b>HR 0.40</b> (0.23 – 0.71)		- 17

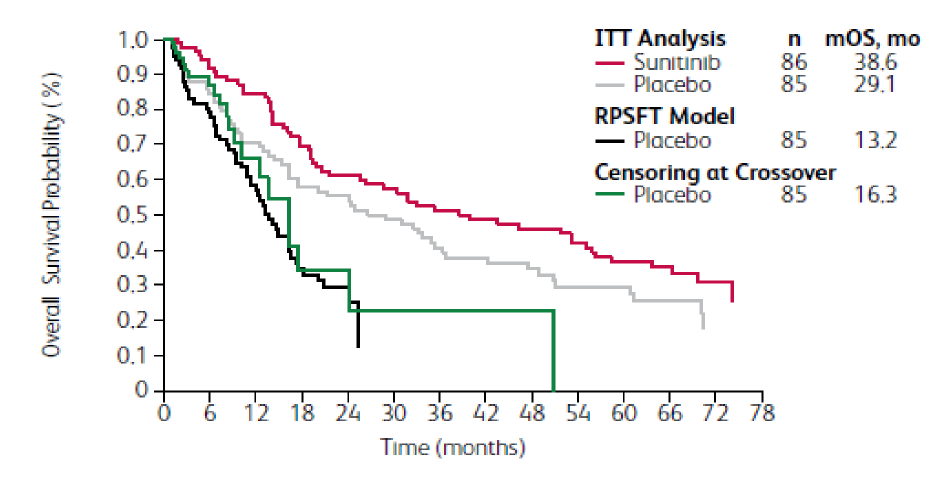
# A6181111 Results (2)

Pfizer submission, section 4.7, pages 42 - 50

	Sunitinib	Placebo
	(N = 86)	(N = 85)
Progression-free survival (F	PFS) ITT population	
Number censored	56	34
Number censored	(65.1%)	(40.0%)
Probability of being event	71.3%	43.2%
free at 6 months (95% CI)	(95% CI, 60.0%, 82.5%)	(30.3%, 56.1%)
Overall-survival (OS) ITT po	pulation	
Namelana	77	64
Number censored	(89.5%)	(75.3%)
Probability of survival at 6	92.6%	85.2%
months (95% CI)	(95% CI: 86.3%, 98.9%)	(95% CI: 77.1%, 93.3%)
<b>Tumour response rates ITT</b>	population (n%)	
Complete recognice (CD)	2	0
Complete response (CR)	(2.3%)	(0.0%)
Partial response (PR)	6	0
	(7.0%)	(0.0%)
Stable/no response (SD)	54	51
Otabierno response (OD)	(62.8%)	(60.0%)

#### A6181111 Overall Survival

Kaplan-Meier estimate of overall survival with and without adjustment for crossover, final analysis, ITT population (source: Raymond et al. 2016<sup>11</sup>)



Source: Pfizer submission, figure 6 (page 48)

# A6181111 subgroup analyses

PFS subgroup analysis (using cox proportional hazards)				
Covariate	Subgroup	N	HR (95% CI)	
Tumour functionality	Functioning	86	<b>0.26</b> (0.13, 0.54)	
rumour functionality	Not Functioning	46	<b>0.75</b> (0.30, 1.84)	
No. of previous	0 or 1	121	<b>0.33</b> (0.19,0.59)	
systemic regimens	≥2	50	<b>0.61</b> (0.27,1.37)	
Previous use of SSA	Yes	68	<b>0.43</b> (0.21,0.89)	
FIEVIOUS USE OF SSA	No	103	<b>0.41</b> (0.22,0.75)	
Sources: Assessment report, table 28, page 96				

### GI and Lung NETs: Clinical Trials

NETTER-1: 177Lu-DOTATATE plus

Trial	RADIANT-4: everolimus plus BSC Vs. placebo plus BSC	NETTER-1: 177Lu-DOTATATE plus octreotide 30mg Vs. octreotide LAR (60 mg)
Design	<ul> <li>Double-blind, randomised, placebo-controlled phase III trial</li> </ul>	<ul> <li>Stratified, open, randomised, comparator-controlled, parallel-group phase III</li> </ul>
Pop	<ul> <li>Patients with advanced, progressive, low- or intermediate- grade GI and Lung NETs</li> </ul>	<ul> <li>Patients with inoperable, progressive (as determined by RECIST Criteria), somatostatin receptor positive, midgut NETs of the small bowel</li> </ul>
Out	<ul> <li>Primary endpoint - PFS (centrally assessed according to RECIST)</li> <li>Secondary endpoints - OS, DoR, ORR and safety</li> </ul>	<ul> <li>Primary endpoint - PFS Independent Review Centre (IRC)</li> <li>Secondary endpoints – OS, DoR, ORR, TTP, safety, tolerability and HRQoL</li> </ul>
Other	<ul> <li>Crossover after progression was not allowed</li> <li>&gt;half previous SSA therapy (mostly for tumour control)</li> <li>Quarter received prior chemo</li> <li>HRQoL: FACT-G questionnaire</li> </ul>	<ul> <li>All patients received prior therapy</li> <li>Concomitant systemic therapy was not permitted</li> </ul>

#### RADIANT-4 results: GI and Lung NETs combined Novartis submission, tables 5.4 and 5.5 (pages 66 - 74)

Progression-free survival (PFS) - central review (Primary data cut, November 2014)

	Everolimus + BSC (n=207)	Placebo + BSC (n=203)
PFS, median, months	<b>11.0</b> (9.2 – 13.3)	<b>3.9</b> (3.6 – 7.4)
HR (95% CI)	<b>0.48</b> (0.35 –	- 0.67)

#### Overall survival (OS) (Secondary data cut, November 2015)

Progressed disease (PD)

OS, median, months	<b>37.16</b> (35.35 – NE)	<b>39.56</b> (23.46 – NE)
HR (95% CI)	0.73 (0.48 -	- 1.11)

rumour response rates (n %) - central review (Primary data cut, November 2014)				
Partial response (PR)	<b>4</b> (2.0)	<b>1</b> (1.0)		
Stable disease (SD)	<b>165</b> (80.5)	<b>62</b> (63.9)		

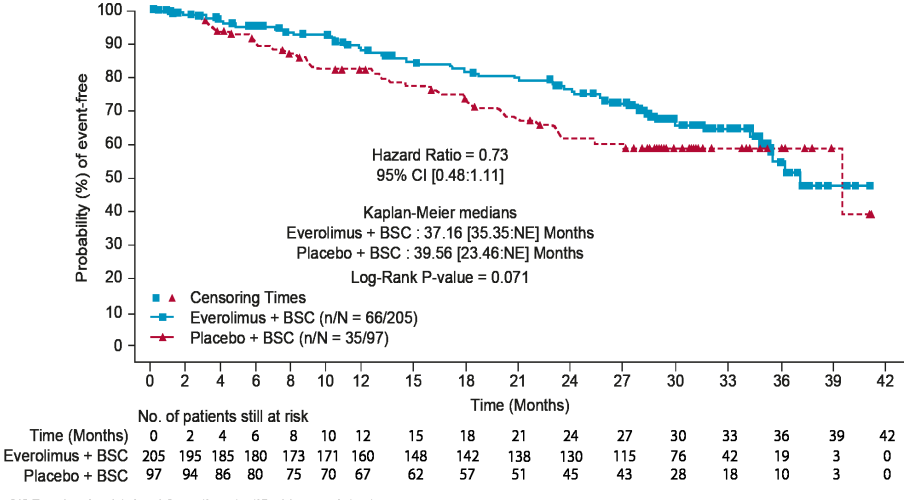
**26** (26.8)

The AG stated that there is little evidence of a difference in PFS within subgroups according to treatment history, previous chemotherapy, previous SSA and tumour grade

**19** (9.3)

#### RADIANT-4 Overall survival: GI and Lung NETs

Kaplan-Meier plot for OS estimates: secondary data cut-off (30th November 2015)



<sup>-[1]</sup> P-value is obtained from the stratified log-rank test.

Source: Novartis submission, figure 5.12, page 73

# RADIANT-4 results: GI NETs only Assessment report, tables 47- 49 (page 110)

Progression-free survival (PFS) - central review				
	Everolimus + BSC (n=118)	Placebo + BSC (n=57)		
PFS, median, months HR (95% CI)	<b>13.1</b> (9.2, 17.3)	<b>5.4</b> (3.6, 9.3)		
	<b>0.56</b> (0.37, 0.84)			
Overall survival (OS)				
OS, median, months HR (95% CI)				
Tumour response rates (n %)				
Stable disease (SD)				
Progressed disease (PD)				

# RADIANT-4 results: Lung NETs only Assessment report, tables 52 - 54 (page 112)

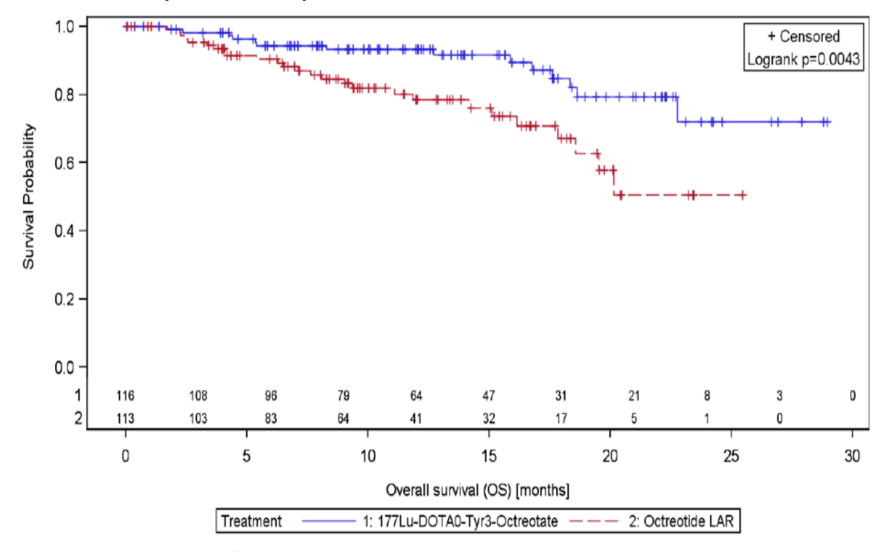
Progression-free survival (PFS)				
	Everolimus + BSC (n=63)	Placebo + BSC (n=27)		
PFS, median, months HR (95% CI)	<b>42</b> (CI not recorded)	18 (CI not recorded)		
	<b>0.50</b> (0.28-0.88)			
Overall survival (OS)				
OS, median, months				
HR (95% CI)				
Tumour response rates (	n %)			
Partial response (PR)				
Stable disease (SD)				
Progressed disease (PD)				

# NETTER-1 Results AAA submission, tables 13 and 14, page 50 –53

Outcomes	Independent IRC			
Progression-free survival (PFS)				
	177 Lu-DOTATATE + Octreotide LAR (n=116)	Octreotide LAR (n=113)		
PFS, median, months	Not reached	8.4		
HR (95% CI)	<b>0.25</b> (0.1	<b>0.25</b> (0.13 – 0.33)		
Patients with events (n)	23	68		
Censored patients (n)	93 45			
Overall survival (OS) (Inter	rim analysis)			
OS, median, months	Not reached	Not reached		
HR (95% CI)	<b>0.398</b> (0.207 – 0.766)			
Patients with events (n)	14	26		
Censored patients (n)	102	87		
Objective response rate (ORR)				
Overall response rate (all patients)	<b>15.5%</b> (10.4 – 25.4)	<b>2.7%</b> (0.0 – 6.3)		

#### **NETTER-1 Overall Survival**

#### OS interim analysis, full analysis set



Source: AAA submission, figure 10, page 52

#### Health-related quality of life (HRQoL)

#### Everolimus

- RADIANT-3
  - Not collected
- RADIANT-4
  - Everolimus had longer median time to definitive deterioration in HRQoL using FACT-G but not statistically significant

#### 177-Lu DOTATATE

- NETTER-1
  - Treatment with 177Lu-DOTATATE does not negatively affect the patient's HRQoL compared with octreotide LAR when using EORTC QLQ-30

#### Sunitinib

- A6181111
  - No statistically significant difference between the sunitinib and placebo groups at any time when using EORTC QLQ-30

#### Adverse events (1)

#### RADIANT-3

- Most common treatment related AEs occurring in ≥20% of patients were rash (52.5%), stomatitis (53.9%), diarrhoea (48%) and fatigue (44.6%)
- 13 incidences of treatment discontinuation due to treatment related AE with everolimus compared to 2 with placebo plus BSC

#### RADIANT-4

- Serious AEs reported for everolimus and BSC were 42.1% and 19.4% respectively
- Most common treatment-related AEs (of any grade) reported by patients receiving everolimus were stomatitis (63%), diarrhoea (31%), fatigue (31%), infections (29%), rash (27%), and peripheral oedema (26%)
- 69 deaths in the trial 20.3% for everolimus and 28.6% for placebo arm

#### Adverse events (2)

#### NETTER-1

- 177Lu-DOTATATE was relatively well tolerated
- Incidence of AE and SAE judged to be treatment-related higher with 177Lu-DOTATATE than with octreotide LAR
- Incidence of grade 3-5 AEs with 177Lu-DOTATATE was low
- Incidences of Grade 3-5 AEs was comparable except for vomiting and lymphopenia

#### A6181111

- AEs more common in the sunitinib group
- Most common treatment-related AEs reported in the sunitinib arm were diarrhoea, nausea and asthenia, all of which were experienced by at least 30% of patients
- Proportion experiencing SAEs was greater in the placebo group (41.5%, versus 26.5% with sunitinib)

### AG's comments on clinical trials

RADIANT-3, RADIANT-4 and A6181111

- All 3 trials double blind low risk of bias in all
- Populations for the 3 trials all in line with the licensed indication for each treatment and with final scope
- Limited information for current prevalence of NETs to assess generalisability
- The applicability of the results in the UK setting was unclear
- High levels of crossover in RADIANT-3 and A6181111 (73% and 69%)

# Company network meta-analyses (P-NETs) Novartis and Pfizer

#### Novartis (everolimus vs sunitinib)

- Bucher indirect comparison using data from RADIANT-3 and A6181111
- PFS local investigator assessment HR 0.83 (0.49, 1.42)
- PFS blinded independent review committee **HR 1.08** (0.59, 1.99)
- OS ITT analysis HR 1.32 (0.81, 2.16)
- OS RPSFT-adjusted analysis HR 1.40 (0.17, 11.72)

#### Pfizer (sunitinib vs everolimus)

 MAIC using patient-level data from A6181111 and aggregate data from RADIANT-3



# Company network meta-analyses

#### **P-NETs**

- Mixed treatment comparison including results from NETTER-1, RADIANT-3 and A6181111
- PFS MTC analysis
  - 177Lu DOTATATE vs everolimus: HR 0.60 (0.04, 9.92)
  - 177Lu DOTATATE vs sunitinib: HR 0.50 (0.03, 8.60)
- OS MTC analysis
  - 177Lu DOTATATE vs everolimus: HR 0.38 (0.07, 2.28)
  - 177Lu DOTATATE vs sunitinib: HR 0.98 (0.15, 6.46)

#### **GINETs**

- Indirect treatment comparison comparing results from NETTER-1 and RADIANT-4
- PFS MTC analysis:
  - 177Lu DOTATATE vs everolimus: HR 0.43 (0.05, 4.24)
- OS MTC analysis:
  - 177Lu DOTATATE vs sunitinib: HR 0.43 (0.09, 2.12)

# AG'S comments on company network metaanalyses (1)

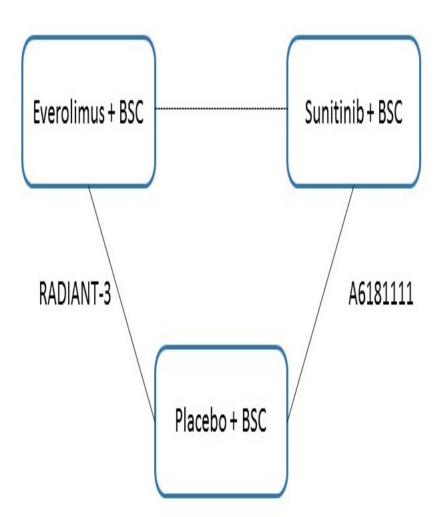
- Novartis submission (Bucher indirect comparison using data from RADIANT-3 and A6181111)
  - Inconsistent results for PFS between central and local review
  - Wide confidence intervals for all results uncertainties
  - Different results when using crossover unadjusted and adjusted results
  - Response rates results with wide confidence intervals suggesting little difference between the two treatments
  - Unclear why Bucher was used over MAIC. However, similar results and Bucher has more mature data
- Pfizer submission (MAIC using patient-level data from A6181111 and aggregate data from RADIANT-3)
  - MAIC here could not adjust for differences in study design across trials
  - RADIANT-3 and A6181111 populations were similar (some differences)
  - Balanced baseline characteristics in RADIANT-3/Imbalanced baseline characteristics in A6181111
  - Small sample size (which after matching halved in size)

### AG'S comments on company network metaanalyses (2)

- AAA P-NETs submission (Mixed treatment comparison using data from NETTER-1, RADIANT-3 and A6181111)
  - No justification that octreotide LAR 60mg is equivalent to placebo, placebo + octreotide (30mg) and placebo + BSC
  - NETTER-1 should be excluded: no patients with P-NETs
  - No consideration of treatment switching for the trials included
  - Wide confidence intervals suggesting uncertainty
  - Models used not reported in the submission and so no comparison of any differences in point estimates
- AAA GI-NETs submission (Indirect treatment comparison comparing results from NETTER-1 and RADIANT-4)
  - No justification that octreotide LAR 60mg is equivalent to placebo, placebo+octreotide (30mg) and placebo+BSC
  - RADIANT-2 should be excluded: population all have functioning tumours (outside MA for everolimus for GI-NETs)
  - For GI NETs populations for OS differ across the studies
  - No consideration of treatment switching for the trials included
  - Wide confidence intervals suggesting uncertainty

# Everolimus Vs Sunitinib (P-NETs)

#### Assessment Group



- RADIANT-3 and A6181111 are comparable to allow an ITC
- Bucher method used but no analyses for heterogeneity between the trials or inconsistency (only 2 trials)
- Outcomes PFS, OS, RR, AEs
- Higher proportion of SSA use in RADIANT-3 (40%) compared to A6181111 (28%),
  - Not thought that this would affect the relative effectiveness of the treatments
- ITC should be interpreted with caution

Source: Assessment report, figure 17 (page 98)

# ITC – PFS results (P-NETs) AG Report

HRs (95% Cls) for (local review) disease progression or death in P-NETs				
Intervention	Comparator	Data source	HR (95% CI)	
Everolimus + BSC	Placebo + BSC	RADIANT-3		
Sunitinib + BSC	Placebo + BSC	A6181111		
Everolimus + BSC	Sunitinib + BSC	Calculated by AG ITC		

Source: Assessment report, table 31 (page 99)

# HRs (95% Cls) for (central review) disease progression or death in P-NETs

Intervention	Comparator	Data source	HR (95% CI)
Everolimus + BSC	Placebo + BSC	RADIANT-3	
Sunitinib + BSC	Placebo + BSC	A6181111	
Everolimus + BSC	Sunitinib + BSC	Calculated by AG	

Source: Assessment report, table 32 (page 99)

# ITC – OS results (P-NETs) AG Report

# HRs (95%CI) for death P-NETs based on final follow-up data (crossover unadjusted)

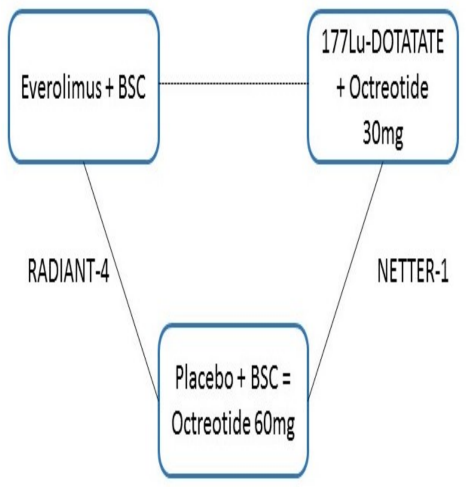
Intervention	Comparator	Data source	HR (95% CI)
Everolimus + BSC	Placebo + BSC	RADIANT-3	
Sunitinib + BSC	Placebo + BSC	A6181111	
Everolimus + BSC	Sunitinib + BSC	Calculated by AG	

HRs (95%CI) for death P-NETs (crossover adjusted RPSFT)				
Intervention	Comparator	Data source	HR (95% CI)	
Everolimus + BSC	Placebo + BSC	RADIANT-3		
Sunitinib + BSC	Placebo + BSC	A6181111		
Everolimus + BSC	Sunitinib + BSC	Calculated by AG		

Sources: Assessment report, table 33 (page 99), Source: Assessment report, table 34 (page 100) and Assessment report, table 35 (page 100)

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# Lutetium-177 Vs everolimus (GI-NETs) Assessment Group



- AG assumed that placebo+BSC can be considered equivalent to octreotide 60mg
- RADIANT-4 includes a combination of GI+lung NETs
- Different tumour locations included under term GI in the two RCTs
  - NETTER-1 only midgut NETs
  - RADIANT-4 fore-, mid- and hindgut NETs
- Bucher used to indirectly compare everolimus to 177Lu-DOTATATE for GINETs: central review PFS, OS, RR and various AEs
- Analyses for heterogeneity or inconsistency between trials was not possible
- ITC should be treated with caution

Source: Assessment report, figure 28 figure 29 (page 141 and 142)

# ITC – PFS & OS results (GI-NETs) AG Report

HRs (95% Cls) for (central review of) disease progression or death in GI NETs				
Intervention	Comparator	Data source	HR (95% CI)	
Everolimus + BSC	Placebo + BSC	RADIANT-4	<b>0.56</b> (0.37, 0.84)	
177Lu-DOTATATE + octreotide 30mg	Octreotide 60mg	NETTER-1	<b>0.21</b> (0.13, 0.33)	
177Lu-DOTATATE + octreotide 30mg	Everolimus + BSC	Calculated by AG ITC	<b>0.37</b> (0.19, 0.69)	

Source: Assessment report, table 67 (page 144)

HRs (95% Cls) for OS in GI NETs				
Intervention	Comparator	Data source	HR (95% CI)	
Everolimus + BSC	Placebo + BSC	RADIANT-4		
177Lu-DOTATATE + octreotide 30mg	Octreotide 60mg	NETTER-1	<b>0.40</b> (0.21, 0.77)	
177Lu-DOTATATE + octreotide 30mg	Everolimus + BSC	Calculated by AG ITC		
Source: Assessment report, table 68 (page 145)				

### Key clinical issues (1)

- The AG's decision problem is in line with the final scope but excluded some comparators/interventions because
  - Interferon alpha not routinely used in practice and no relevant studies
  - Chemotherapy no relevant study to include in the network
  - Lutetium (P-NETs) population not included in NETTER-1
  - AAA presented MTC and economic analysis for P-NETs using data from NETTER-1, which the AG considered inappropriate
     Have the appropriate comparisons been made for each tumour locations?
- What conclusions can be drawn from the ITC for P-NETs, given:
  - Exclusion of lutetium from the network
  - High-level of crossover in RADIANT-3 and A618111, RPSFT-adjusted results also presented
  - AG considered the RADIANT-3 and A6181111 to be comparable, although they differed in SSA use – 40% vs 28% respectively
    - not considered by AG to affect the relative effect of the treatments
  - A6181111 included both functioning and non-functioning tumours, but the secretory profile in RADIANT-3 was not reported
  - Results showed
    - Can they be assumed to be clinically equivalent?

### Key clinical issues (2)

- What conclusions can be drawn from the ITC for GI NETs, given:
  - The assumption that 60mg octreotide is clinically similar to placebo + BSC?
  - AG ITC used the full population from NETTER-1 and a subset of RADIANT-4 (GI only)
    - AAA used the full RADIANT-4 population (GI and lungs)
  - Comparability of RADIANT-4 and NETTER-1
    - GI (fore-, mid- and hind-gut) vs midgut NETs respectively
    - Non-functioning vs mixed (functioning and non-functioning)
    - All patients in NETTER-1 were somatostatin receptor positive, but not known for RADIANT-4
  - The inclusion of RADIANT-2 by AAA
    - Excluded in the AG ITC because the population is outside the MA for everolimus