# NATIONAL INSTITUTE FOR HEALTH AND CLINICAL EXCELLENCE

## Single Technology Appraisal (STA)

### Everolimus for the second-line treatment of metastatic renal cell carcinoma

The following documents are made available to consultees and commentators:

- Response to consultee, commentator and public comments on the Appraisal Consultation Document (ACD) prepared by the Technical Lead, Helen Tucker
- Manufacturer's updated cost-effectiveness analysis in response to the final appraisal determination (confidential information removed)
  - o Additional sensitivity analysis
  - o Email from manufacturer reporting the mean ICERs
  - o Amended PAS submission
- ERG summary response to manufacturer submission prepared by Peninsula Technology Assessment Group (PenTAG) (confidential information removed)
- ERG response to Additional Sensitivity Analyses prepared by Peninsula Technology Assessment Group (PenTAG) (confidential information removed)

#### NATIONAL INSTITUTE FOR HEALTH AND CLINICAL EXCELLENCE

Everolimus for the second-line treatment of advanced renal cell carcinoma

Response to consultee, commentator and public comments on the Appraisal Consultation Document (ACD)

### **Definitions:**

**Consultees –** Organisations that accept an invitation to participate in the appraisal including the manufacturer or sponsor of the technology, national professional organisations, national patient organisations, the Department of Health and the Welsh Assembly Government and relevant NHS organisations in England. Consultee organisations are invited to submit evidence and/or statements and respond to consultations. They are also have right to appeal against the Final Appraisal Determination (FAD). Consultee organisations representing patients/carers and professionals can nominate clinical specialists and patient experts to present their personal views to the Appraisal Committee.

**Clinical specialists and patient experts** – Nominated specialists/experts have the opportunity to make comments on the ACD separately from the organisations that nominated them. They do not have the right of appeal against the FAD other than through the nominating organisation.

**Commentators** – Organisations that engage in the appraisal process but that are not asked to prepare an evidence submission or statement. They are invited to respond to consultations but, unlike consultees, they do not have the right of appeal against the FAD. These organisations include manufacturers of comparator technologies, NHS Quality Improvement Scotland, the relevant National Collaborating Centre (a group commissioned by the Institute to develop clinical guidelines), other related research groups where appropriate (for example, the Medical Research Council and National Cancer Research Institute); other groups (for example, the NHS Confederation, NHS Information Authority and NHS Purchasing and Supplies Agency, and the *British National Formulary*).

**Public** – Members of the public have the opportunity to comment on the ACD when it is posted on the Institute's web site 5 days after it is sent to consultees and commentators. These comments are usually presented to the appraisal committee in full, but may be summarised by the Institute secretariat – for example when many letters, emails and web site comments are received and recurring themes can be identified.

## Comments received from consultees

Consultee	Comment	Response
The James Whale Fund for Kidney Cancer	Whether we consider that all the relevant evidence has been taken into account. It is our assertion that meaningful patient input is missing from the ACD. The James Whale Fund feel the evidence should be revisited and the patient perspective must be included and given due weight if N I C E wish to present a balanced and rounded appraisal.	The patient perspective was acknowledged by the Committee. See FAD sections 4.1, 4.2 and 4.3
The James Whale Fund for Kidney Cancer	The spend on cancer drugs is higher in other EU Countries. A recent report from Policy Exchange states that spending on cancer medicines in England is only 60% of that spent by other advanced EU countries and our cancer death rate is 6% higher than the EU average, it would be naïve not to see the connection between those two figures. Cancer patients in England are hugely disadvantaged by this process of rationing by cost.	Comment noted. The Institute recognises that guidance from other organisations may differ from its own guidance, because of different criteria for making decisions. The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the manufacturer's submission and the ERG report.
The James Whale Fund for Kidney Cancer	The last 10 years has seen much research into innovative anti-cancer drugs come to fruition. In the case of Kidney Cancer, NICE has reviewed 5 such new drugs and has only approved one 1 <sup>st</sup> line new drug (Sunitinib) and refused all 2 <sup>nd</sup> line sequential treatments. The drugs refused by N I C E are widely available in all western countries and NICE's justification for denying access to innovative new cancer drugs to NHS patients are based on esoteric cost calculations and statistics which are incomprehensible to patients and the general public. Denying treatment to terminally ill cancer patients has been hugely controversial and the Department of Health, through N I C E, has been forced to react to public criticism by introducing an "End of Life" criteria to ensure that modern and comparably costly drugs, are not automatically refused when they fail the notorious and arbitrary N I C E QALY. There is no evidence that the EOL criteria have been applied to this application for Everolimus even though Everolimus fits the criteria perfectly. The consequence of this unfair approach is that mRCC patients have only 1 drug for 1 <sup>st</sup> line treatment (accepting there maybe some limited use for 20 year old immunotherapy treatments such as interferon alfa), none at all for sequential 2 <sup>nd</sup>	Comments noted. The Appraisal Committee considered the supplementary advice and it agreed that everolimus does fulfil the criteria as a life extending, end-of-life treatment. However, the Committee concluded that everolimus for the second-line treatment of advanced RCC would not be a cost-effective use of NHS resources. See FAD sections 4.14 and 4.15.

Consultee	Comment	Response
	line treatment leaving only, as a last resort, best supportive care. Once again kidney cancer patients in the UK are disadvantaged by the N I C E model of cost analysis.	
The James Whale Fund for Kidney Cancer	The figure of the £30,000 Q A L Y has not been updated since its inception – one can imagine the furore if other cost areas in the NHS i.e. salaries and expenses had remained unchanged for 9 years. A simple calculation shows if the QALY had been adjusted in line with other NHS costs, a £50/55,000 Q A L Y would be the norm and taking the figure of 1.4 quoted recently by Professor Stevens as the multiplier, the EOL Q A L Y should now be £70/75,000. N I C E appears to exist in a time warp for this one area of their work. Today's treatments for today's patients should not be judged against a set of "rules" which are nearly 10 years old.	Comment noted
The James Whale Fund for Kidney Cancer	(patient quote) "Cancer survival rates are much higher in other EU countries especially when sequential treatment is available."	Comment noted. The Institute recognises that guidance from other organisations may differ from its own guidance, because of different criteria for making decisions. The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the manufacturer's submission and the ERG report.
The James Whale Fund for Kidney Cancer	Whether we consider that the summaries of the clinical effectiveness and cost effectiveness are reasonable interpretations of the evidence and the preliminary views on the resource impact and implications for the NHS are appropriate         It is apparent to us from talking and listening to patients and the general public that the majority of people do not understand the pseudo-science of mathematical models,         ICER's and OALX's Deticate do not understand how an actual implication set of C21 000 per set of C	Comment noted The reference case stipulates that the cost effectiveness of treatments should be expressed in terms of incremental cost per quality- adjusted life year. See Guide to the Methods of Technology Appraisal
	ICER's and QALY's. Patients do not understand how an actual invoice cost of £31,000 pa can, following an appraisal by N I C E, be transformed into a cost to the NHS of £75,000 pa. If N I C E cannot find a way to explain their processes to patients denied access to	section 5.2.11 (Available from URL http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p

Consultee	Comment	Response	
	clinically effective treatments that Clinicians wish to prescribe, then we suggest it is out of touch with the NHS patients it is meant to be serving.	df)	
The James Whale Fund for Kidney Cancer	patient quote) "It's so difficult to understand what they are saying, with all that gobblygook, when Sutent stops working for me, can I really expect to live another 11 or 12 months without any proper cancer treatment at all. That's not what I read on the patients forums. Do other stage 4 patients and the Oncologists agree with that I wonder?"	need and that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of the technologies (Social Value Judgments - Principles for the development of NICE guidance; principle 5).	
The James Whale Fund for Kidney Cancer	NICE should take into account the wider societal benefits of access to end of life drugs for cancer patients when assessing cost effectiveness. If patients on active treatment can continue to work and support their families, is that worth nothing?	Comment noted The reference case stipulates that that, the perspective on outcomes should be all direct health effects, whether for patients or, when relevant, other people (principally carers). The perspective adopted on costs should be that of the NHS and PSS. See Guide to the Methods of Technology Appraisal sections 5.2.7 and 5.5.10 (Available from URL http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p df)	
The James Whale Fund for Kidney Cancer	(patient quote) " The NHS has a forecast underspend against budget this year of £1.4 billion – is it a cost effective use of NHS resources to keep that money sitting in NHS bank accounts rather than spend it on front line services like cancer treatments for patients who desperately need them."	Comment noted.	

Consultee	Comment	Response
The James Whale Fund for Kidney Cancer	If this decision is not changed , NICE will have recently rejected all five 2 <sup>nd</sup> line kidney cancer treatments despite promised greater flexibility from NICE for EOL drugs	Comment noted. The Committee noted that there is an unmet clinical need and that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of the technologies (Social Value Judgments - Principles for the development of NICE guidance; principle 5).
The James Whale Fund for Kidney Cancer	Is there a figure being used as the benchmark for "end of life" drugs? How do patients or the public know whether that figure is "reasonable"? How can we comment when the information is not made available? What is a cost effective use of resources when keeping any patient alive? Is it the cost of kidney dialysis per year; is it the cost of an organ transplant operation and ongoing drugs for life?	The supplementary advice does not suggest for Committee to apply a particular weight for the cost effectiveness estimate to fall within the acceptable threshold range. The Committee is asked to come to a value judgment on whether the magnitude of additional weight, that would need to be assigned to the original QALY benefits in the patient group for the cost effectiveness of the drug to fall within the current threshold range, would be acceptable in light of the evidence presented.
The James Whale Fund for Kidney Cancer	(Patient quote)"Our drugs will always be more expensive as there are far fewer of us and pharmaceutical companies have to recoup R & D costs. Drugs must cost the same to get a license whether they are prescribed to 1000 rarer cancer patients or 40,000 patients."	Comment noted. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL

Consultee	Comment	Response
		http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p df)
The James Whale Fund for Kidney Cancer	(patient quote) "Everolimus is cost effective – it works, it does what it says on the tin. I know what it is worth because I'm taking the drug."	Comment noted. The Committee noted that there is an unmet clinical need and that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of the technologies (Social Value Judgments - Principles for the development of NICE guidance; principle 5).
The James Whale Fund for Kidney Cancer	(patient quote) "N I C E is just rationing treatments based on money, but rarer cancer patients obviously are still coming off worse".	Comment noted. See response above, in addition, the Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p df)
The James Whale Fund for Kidney Cancer	Whether we consider that the provisional recommendations of the Appraisal Committee are sound and constitute a suitable basis for the preparation of guidance to the NHS.	Comment noted. The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the
	The general feeling from the kidney cancer community is that they are passionate defenders of the NHS and the principle of universal care, but do not understand why a committee set up to appraise cancer drugs would do so without a leading Oncologist on	manufacturers' submission and the ERG report. The Committee noted that everolimus was likely to be

Consultee	Comment	Response
	the panel and without the added value and experience of a cancer patient. To exclude both viewpoints from membership of the Appraisal committee in favor of multiple commissioning and health economics input seems perverse.	clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.
The James Whale Fund for Kidney Cancer	(patient quote) "Rarer cancer patients are discriminated against & feel disenfranchised by the NICE process"	Comment noted. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p df)
The James Whale Fund for Kidney Cancer	patient quote) "Kidney cancer patients have paid into the NHS ; I've paid a lifetime of taxes - we have paid into the system now all we want is to have treatment options like other cancer patients"	Comment noted. In developing clinical guidance for the NHS, no priority should be given based on individuals' income, social class or position in life and individuals' social roles, at different ages, when considering cost effectiveness (SVJ principle 8).
The James Whale Fund for Kidney Cancer	(patient quote) "This QALY figure is arbitrary, it is out of date and based on goodness knows what? Was it guesswork?"	Comment noted The reference case stipulates that the cost effectiveness of treatments should be expressed in terms of incremental cost per quality-

Consultee	Comment	Response
		adjusted life year. See Guide to the Methods of Technology Appraisal section 5.2.11 (Available from URL http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p df)
The James Whale Fund for Kidney Cancer	<ul> <li>The James Whale Fund for Kidney Cancer ask the Appraisal Committee to take account of the following general points from the perspective of the hundreds of kidney cancer patients who will be affected by their ultimate decision.</li> <li>We feel the principle of cost effectiveness is applied randomly – N I C E asserts it is the guardian of NHS resources by applying clinical effective evidence in a rigorous manner. It tells us that NHS funded treatments must be evidence –based. Despite this assertion cancer patients know there is striking evidence this principle is not consistant across the NHS. It is difficult for kidney cancer patients to reconcile the control N I C E exerts over clinically effective and proven cancer drugs and yet fails to apply to other NHS funded treatments –</li> <li>1. Homeopathy, which is available on the NHS at huge cost and yet is unproven and felt by many to be no better than placebo.</li> <li>2. Acupuncture, which is available on the NHS with very little peer reviewed evidence.</li> <li>3. Alternative medicines available on the NHS and not subject to NICE scrutiny.</li> <li>4. The swine flu panic now agreed to have led to the waste of huge NHS resources 5. The winter flu jab for the over 65's, now seen as failing to deliver measurable benefit.</li> </ul>	Comments noted. The Committee does not consider the affordability that is costs alone, of new technologies but rather their cost effectiveness in terms of how its advice may enable the more efficient use of available
	These examples are proof to patients the NHS is not consistent and N I C E is a questionable guardian of precious NHS resources and yet N I C E persist in denying treatments to fulfil an unmet clinical need for a 2 <sup>nd</sup> line treatment for terminally ill kidney cancer patients.	

Consultee	Comment	Response
The James	Patients tell us they are actively encouraged to enter clinical trials for new cancer drugs.	Comment noted
Whale Fund	They do so for a number of reasons; it may be the only route to active treatment, they	
for Kidney	feel they are "doing a good thing" helping to further medical knowledge and they feel their	
Cancer	involvement may help future generations of cancer patients. Each time that N I C E deny	
	access to effective drugs, the effect on those patients who took part on the clinical trials is	
	immediate and diminishes their contribution; they feel let down and some feel	
	hoodwinked. Their hopes of enabling effective treatment to be used to help other cancer	
	patients are dashed. The knock on effect for further research and trials in the UK must be	
	recognized as must the effect on patients whose hopes are raised when they hear first	
	hand in their Clinics, about good results and evidence, but then discover N I C E will not	
	allow these new compounds to be funded by the NHS.	
The James	We urge the committee on the 9 <sup>th</sup> March to acknowledge the value of the patient	Comment noted. The Committee
Whale Fund	experience, we have asked that our expert patient should be available for	considered all the evidence submitted,
for Kidney	your committee to talk to about the points we have raised in our submission and we	including evidence from clinical trials,
Cancer	would like your agreement to that request.	patient and clinical experts, the
		manufacturers' submission and the
	In conclusion we will share with your committee the words of a stage 4 kidney cancer	ERG report. The Committee noted
	patient who, until disease progression 3 months ago, was taking a kidney cancer drug	that everolimus was likely to be
	refused by N I C E, a cancer drug that has given him 3 years of extra life - not a few	clinically effective. However, for both
	weeks as we hear quoted in the media, but 3 years during which time he has continued	legal and bioethical reasons the
	to work and play a full role in his family	Committee must take account
	"Being told you have terminal kidney cancer is not the worst thing in the world to	economic considerations and cost
	happen to you – far worse is knowing there are proven drugs that can help you,	effectiveness of everolimus (Social
	but you can't have them."	Value Judgments - Principles for the
	Detients in this situation new need convertice Ord line treatments when is using to sit this	development of NICE guidance;
	Patients in this situation now need sequential 2nd line treatment: who is going to sit this	principle 5). The Committee
	patient down and say to him	concluded that everolimus would not
	"It has become too expensive for we to keep you alive "	be a cost-effective uses of NHS
	"It has become too expensive for us to keep you alive."	resources.

Consultee	Comment	Response
The James Whale Fund for Kidney Cancer	Are there any equality related issues that need special consideration that are not covered in the ACD? Do kidney cancer patients just have the "wrong type of cancer" Patients are dying prematurely because they simply have the bad luck to have been diagnosed with a rare cancer, through no fault of their own. Nothing will change until the NHS accepts that rarer cancer patients need a separate process of appraisal. A one size HTA does not fit all.	Comment noted. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p df)
The James Whale Fund for Kidney Cancer	patient quote) "Everolimus is available in other EU countries as 2 <sup>nd</sup> line treatment for mRCC, why not in Great Britain?"	Comment noted. The Institute recognises that guidance from other organisations may differ from its own guidance, because of different criteria for making decisions. The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the manufacturer's submission and the ERG report.
The James Whale Fund for Kidney Cancer	(patient quote) "KC patients have limited treatment options unlike more common cancers ( chemotherapy & radiotherapy do not work for kidney cancer) Why can't similar amounts of money that other cancer patients have access to for their treatments be given to us to help pay for drugs we need. If patients with rarer cancers can't get treatment because they are in a minority surely this is a form of discrimination."	Comment noted. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p df)
The James Whale Fund for Kidney	(patient quote) "The majority of KC patients are aged 60+; not everyone has access to computers and NICE website is awful, it is not user friendly, it puts you off before you start; how we are expected to appeal properly. We only have 20 days to appeal against	Comment noted

Consultee	Comment	Response
Cancer	refusal for our drugs and yet this was referred to NICE in November 2008. The NHS and the N I C E Quango take as much time & money as they want to get their arguments marshalled, but again we get no help, no resources at all to put our case forward."	
The James Whale Fund for Kidney Cancer	(patient quote) "The Human Rights Act, article two, gives every human being THE RIGHT TO LIFE, denial of a proven clinically effective treatment which gives an individual that right cannot therefore be legal under the convention."	Comment noted. The Appraisal Committee has been given supplementary advice to be taken into account when appraising treatments which may be life extending. This advice addresses the notion of additional benefits not readily captured in the reference case and to have regard to the importance of supporting the development of innovative treatments that are (anticipated to be) licensed for small groups of patients who have an incurable illness. The Committee concluded that everolimus fulfilled the end-of-life criteria and considered it as such.
Kidney Cancer UK	Do you consider that all of the relevant evidence has been taken into account? Not in our view.	The patient perspective was acknowledged by the Committee. See FAD sections 4.1, 4.2 and 4.3.
	Evidence on patient benefits has scarcely been considered in the ACD, compared with the enormous amount of space devoted to discussion of the evidence on costs. In our view the central measure of a QALY is a woefully inadequate measure of patient benefit, calibrated as it is on the basis of a number of truly heroic assumptions. Patient benefit encompasses far more than a QALY.	The reference case stipulates that the cost effectiveness of treatments should be expressed in terms of incremental cost per quality-adjusted life year. See Guide to the Methods of Technology Appraisal section 5.2.11

Consultee	Comment	Response
		(Available from URL http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p df)
Kidney Cancer UK	A more academically respectable approach to the evaluation would have involved calculation of net present values [NPVs] in a full-blown cost-benefit analysis. Admittedly, NPV calculations would be much more difficult to make, given that they would require direct valuation of patient benefits. But in this-as in everything else-there is more to be said for <i>rough</i> estimates of the <i>precise</i> concept than for <i>precise</i> estimates of some <i>rough</i> concept. An incremental cost effectiveness ratio [ICER] per QALY is a pretty rough concept; and sometimes it is, solemnly, and most precisely, given down to the last £1.	Comment noted. The reference case stipulates that the cost effectiveness of treatments should be expressed in terms of incremental cost per quality- adjusted life year. See Guide to the Methods of Technology Appraisal section 5.2.11 (Available from URL http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p df
Kidney Cancer UK	Do you consider that the summaries of clinical and cost effectiveness are reasonable interpretations of the evidence, and that the preliminary views on the resource impact and implications for the NHS are appropriate? Not in our view The summaries rest very heavily on certain assumptions regarding how long patients can survive solely on best supportive care after treatment with Sunitinib has failed. PenTAG's ICER per QALY of £75,000 is associated with a mean survival of 11 months in the best-supportive- care arm. But there are reasons to believe that 11 months is an unrealistic estimate of survival on best supportive care. For instance, in a paper by Di Lorenzo et alia published in <i>The Journal of Clinical Oncology</i> [10.1200/JCO, 2009, August] it is shown that patients failing on Sunitinib and then going on to receive Sorafenib as second-line treatment lived for a median period of just 32 weeks [or a little less than 7.4 months]. It seems inconceivable that patients on best supportive care would survive longer than patients receiving an active drug.	The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the manufacturers' submission, revised cost-effectiveness estimates submitted in response to the appraisal consultation document and corresponding ERG reports. It also carefully considered the comments received from consultees and commentators in response to the Appraisal Consultation Document. The Committee noted that the overall survival associated with best supportive care in the ERG analysis

Consultee	Comment	Response
		(10.8 months) was likely to be higher
		than in clinical practice. The
		Committee noted the difference in
		overall survival between treatment
		arms was 8.2 months in the
		manufacturer's revised RPSFT
		analysis and 5.9 months in the ERG's
		revised RPSFT analysis. It noted
		evidence from the manufacturer and
		the clinical specialists that an increase
		in overall survival of 1.4 months per
		month of increased progression-free
		survival would be considered
		plausible. Therefore, the Committee
		agreed that the incremental overall
		survival derived using the
		manufacturer's revised RPSFT
		analyses was almost twice as much
		as would be expected, given that the
		trial had observed an increase of 3
		months in progression-free survival.
		The Committee accepted that the
		ERG's estimate of the incremental
		difference in overall survival for
		everolimus versus best supportive
		care (5.9 months) was more plausible
		than that derived by the manufacturer
		and was based on all the available
		data. See FAD section 4.10

Consultee	Comment	Response
Kidney Cancer UK	A further piece of evidence is found in a study by Z. Liu et alia presented at the Joint 15 <sup>th</sup> Congress of the European CanCer Organisation [ECCO] and 34 <sup>th</sup> Congress of the European Society for Medical Oncology [ESMO] Berlin, 20-24 September 2009. In this study, the median overall survival for patients who received no active treatment after Sunitinib is found to be only 5.2 months.	See above response.
Kidney Cancer UK	Do you consider that the provisional recommendations of the Appraisal Committee are sound and constitute a suitable basis for the preparation of guidance to the NHS? Not in our view We feel that, on more realistic assumptions regarding relative survival, the ICER per QALY for Everolimus would come down to around the same level as that at which Sunitinib was approved for NHS funding, namely £54,000. We note that it is accepted that, like Sunitinib, Everolimus is deemed eligible to be designated as an end-of-life medicine. Accordingly, we suggest that the final decision on Everolimus be aligned with that on Sunitinib.	The supplementary advice does not suggest for Committee to apply a particular weight for the cost effectiveness estimate to fall within the acceptable threshold range. The Committee is asked to come to a value judgment on whether the magnitude of additional weight, that would need to be assigned to the original QALY benefits in the patient group for the cost effectiveness of the drug to fall within the current threshold range, would be acceptable in light of the evidence presented.
Royal College of Nursing	<b>Do you consider that all of the relevant evidence has been taken into account?</b> We are unaware of any evidence that has not been included in this technology appraisal	Comment noted. No action required.
Royal College of Nursing	Do you consider that the summaries of clinical and cost effectiveness are reasonable interpretations of the evidence, and that the preliminary views on the resource impact and implications for the NHS are appropriate? We agree with the interpretations of the clinical evidence. We do not have enough expertise to comment on cost- effectiveness and the methodology used.	Comment noted. No action required.

Consultee	Comment	Response
Royal College of Nursing	Do you consider that the provisional recommendations of the Appraisal Committee are sound and constitute a suitable basis for the preparation of guidance to the NHS? It is regretful that the preliminary recommendations contained in the document, mean that	Comment noted. No action required.
	a second line treatment would not be available to patients but note that these	
	recommendations are in line with the previous technology appraisal of Sorafenib.	
Royal College of Nursing	Are there any equality related issues that need special consideration that are not covered in the ACD? We are not aware of any equality related issues that need special consideration which	Comment noted. No action required.
	have not been covered in the ACD.	
Royal College of Physicians	I write on behalf of the NCRI/RCP/RCR/ACP/JCCO with relation to this ACD consultation. We are grateful for the opportunity to respond and would like to make the following comments. Our thanks go to our clinical expert nominee, for coordinating the response. We are disappointed with the ACD decision not to fund everolimus for second line treatment of patients with metastatic renal cancer, after failure of sunitinib therapy. The evidence review group agreed that everolimus has been shown to be clinically effective, increasing survival by 3 months, and meets the end-of-life criteria for drug funding. However, they have declined funding for this small group of patients purely on the basis of cost. Health economic analyses are sensitive to small changes in inputted data and there is often disagreement, even amongst the experts, about interpretation of the results.	Comment noted. The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the manufacturers' submission and the ERG report. The Committee noted that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.

Consultee	Comment	Response
Royal College of Physicians	We are concerned that the committee have misunderstood the prognosis for this group of patients. Section 4.15 quotes 'life expectancy for people with advanced RCC receiving best supportive care alone was unlikely to be greater than 24 months and was potentially as low as 6 months'; this range is actually for patients receiving first line sunitinib. For patients who fail sunitinib, the likely survival without further active treatment is only in the region of 4 or 5 months (expert opinion). It is important that the committee reconsider their decision in the light of this misunderstanding.	Comment noted. This has been amended in the FAD. See section 4.15.
Royal College of Physicians	It is also important to note that the results of the RECORD-1 clinical trial. Patients in both arms of the study received further lines of therapy after everolimus, resulting in a median overall survival of 13 months. This would not of course be the case for the British public with advanced renal cancer. They will be able to receive sunitinib (now NICE approved) but no further treatment if this ACD is ratified in the Final Appraisal Determination.	Comment noted. The Committee understood that there had been crossover after disease progression and that statistical techniques to control for this crossover were necessary. See FAD section 4.6. The Committee noted that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.
Expert 1	The rejection of Everolimus in the context of NICE's rejection of avastin, nexavar and torisel means that oncologists have only 1 drug –sutent –to treat advanced RCC in the	Comment noted. The Appraisal Committee has been given
	first line and no second line treatments with the exception of old discredited drugs such as interleukin or interferon. This situation means that patients are denied modern drugs	supplementary advice to be taken into account when appraising treatments

Consultee	Comment	Response
	which prolong life simply on cost grounds. This is neither moral nor just.	which may be life extending. This advice addresses the notion of additional benefits not readily captured in the reference case and to have regard to the importance of supporting the development of innovative treatments that are (anticipated to be) licensed for small groups of patients who have an incurable illness. The Committee concluded that everolimus fulfilled the end-of-life criteria and considered it as such.
Expert 1	Everolimus fits the EOL criteria –short like expectancy, 3 months plus life extension, ICER over £30k per annum, no available alternatives I can see no evidence that NICE have taken these criteria into account . They were designed specifically to cope with the problems encountered by very expensive life prolonging drugs and yet they have been ignored.	Comments noted. The Committee took in to account the end of life criteria in reaching its decision. See FAD sections 4.14 and 4.15
Expert 1	RCC patients are being discriminated against by the nature of the QALY which turns a cost of £ 30000 into a fantastic figure of over £ 70000 per annum. The application of the current rule set and methodology means that it is almost certain that all modern new drugs for RCC will be rejected leaving England in a situation where these life extending drugs are denied whereas they are widely available across Europe and the USA	Comment noted. The reference case stipulates that the cost effectiveness of treatments should be expressed in terms of incremental cost per quality- adjusted life year. See Guide to the Methods of Technology Appraisal section 5.2.11 (Available from URL http://www.nice.org.uk/media/B52/A7/ TAMethodsGuideUpdatedJune2008.p df)
Expert 1	There is no doubt that Everolimus is clinically effective in extending life. RCC is a lethal disease with very poor outcomes. NICE has placed patients in a situation where life extending drugs have been denied on the grounds of cost and cost alone but a cost	Comment noted. The Committee agreed that everolimus was likely to be clinically effective. However, for

Consultee	Comment	Response
	based on the strange world of the QALY which no patient or carer can understand and is unrecognisable in the real world. Patients deserve transparency and not to be at the mercy of cold blooded health economics.	both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.
Novartis	Everolimus is licensed for the treatment of patients with advanced renal cell carcinoma (aRCC), whose disease has progressed on or after treatment with VEGF-targeted therapy. The only current NICE approved therapy for 1 <sup>st</sup> -line treatment of aRCC is the VEGF-targeted therapy sunitinib. Therefore in the absence of everolimus there are no other effective treatment options available for UK patients via the NHS.	Comment noted. The Committee heard from clinical specialists and patient experts that there are limited treatment options for people with advanced RCC. See FAD section 4.2
Novartis	The preliminary decision to not recommend everolimus is based on the estimates of cost- effectiveness presented by PenTAG. It was felt that Novartis had under-estimated overall survival (OS) in the best supportive care (BSC) arm using both modelling approaches presented. In order to correct for this perceived underestimation of OS in the BSC arm in the Novartis models, PenTAG made various adjustments which resulted in incremental cost-effectiveness ratios (ICERs) of £65,200 and £75,700 (IPCW and RPSFT methods respectively). However, as the difference in OS between everolimus and BSC is one of the biggest influences on the resulting incremental cost-effectiveness ratio (ICER) it is important that the estimates of OS in the BSC arm are realistic and justified based on the available evidence.	Comment noted. The Committee noted that the overall survival associated with best supportive care in the ERG analysis (10.8 months) was likely to be higher than in clinical practice. The Committee noted that the difference in overall survival between treatment arms was 8.2 months in the manufacturer's revised RPSFT analysis and 5.9 months in the ERG's revised RPSFT analysis. It noted evidence from the manufacturer and the clinical specialists that an increase in overall survival of 1.4 months per month of increased progression-free survival would be

Consultee	Comment	Response
		considered plausible. Therefore, the Committee agreed that the incremental overall survival derived using the manufacturer's revised RPSFT analyses was almost twice as much as would be expected, given that the trial had observed an increase of 3 months in progression-free survival. The Committee accepted that the ERG's estimate of the incremental difference in overall survival for everolimus versus best supportive care (5.9 months) was more plausible than that derived by the manufacturer and was based on all the available data. See FAD section 4.10
Novartis	Critically, it is noted that the ICER of £75,700 presented by PenTAG is based on an OS in the BSC arm of 10.9 months (discounted). Novartis strongly believe a mean OS of 10.9 months is not reflective of clinical outcomes in patients who fail 1 <sup>st</sup> -line sunitinib therapy and then receive only BSC in the 2 <sup>nd</sup> -line setting. As with any modelling if the results are not reflective of clinical reality then the resulting ICERs need to be challenged.	Comment noted. See above response.
Novartis	In the original submission Novartis presented an economic analysis based on the Inverse Probability Censoring Weights (IPCW) statistical approach using the February 2008 data cut of the pivotal, phase III, everolimus, RECORD-1 trial. In response to comments in the ERG Report that a rank preserving structural failure time (RPSFT) statistical approach might be preferable, Novartis undertook to conduct the RPSFT analysis, based on the November 2008 data cut, and presented the results within a two week timeframe. This was conducted in the hope of providing a more comprehensive evidence base to inform the Appraisal Committee and thus facilitate a faster decision. Both the IPCW and RPSFT economic analyses presented by Novartis were subsequently adjusted by PenTAG to	The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the manufacturers' submission, revised cost-effectiveness estimates submitted in response to the appraisal consultation document and corresponding ERG reports. It also

Consultee	Comment	Response
	allow for a perceived underestimation of survival in the BSC arm of the economic model. However, based on an additional body of evidence described below, including the views of national and international clinical experts experienced in treating aRCC, the PenTAG adjustment to the RPSFT analysis giving a cost/QALY of £75,700 is not clinically plausible as it relies on an estimate of mean survival in the BSC arm of 10.9 months (11.2 months undiscounted). Moreover we have been able to show by statistical means that the PenTAG suggested correction to the IPCW model results in an overall, effective hazard ratio of 0.6 rather than the intended HR of 0.55. This means that survival in the BSC arm is over-estimated thus inflating the ICER. In order to provide the Appraisal Committee with the most plausible and robust estimates of cost-effectiveness we have updated both of our analyses (IPCW and RPSFT) to take into account PenTAG's criticisms and incorporate the longest term clinical data from the RECORD-1 trial ie data from the November 2008 analysis. This has resulted in revised estimates of cost- effectiveness of £49,537/QALY (RPSFT) and £52,648/QALY (IPCW). The underlying estimates of mean overall survival in the BSC arm are 7.9 months and 9.6 months respectively (discounted values).	carefully considered the comments received from consultees and commentators in response to the Appraisal Consultation Document.
Novartis	Based on the evidence that we have compiled since the ACD was issued, the latter estimates of survival are more plausible, than the estimates presented by PenTAG. These updated incremental cost-effectiveness ratios are lower than those previously presented due to the greater survival demonstrated in the longer term November 2008 analysis and the addition of further cycles in the model to capture the additional benefit. The original model was developed for the February 2008 data cut and therefore only required 18 cycles to capture the available data. However, as stated in our submission of the RPSFT analysis, due to the fact that the November 2008 data-cut suggests greater survival than the February 2008 data-cut, there are more everolimus patients still alive in the final cycle (cycle 18) of the original economic model. Unfortunately, there was insufficient time to add further cycles to the model to account for this when we submitted the RPSFT analysis (due to the 2 week turnaround required) but we have now been able to update the model in order that all of the benefits of everolimus can be reflected in the economic analysis [39 cycles are required to fully account for the additional survival]. The overall impact of allowing this greater survival to be taken into account in the model has	The Committee discussed the manufacturer's revised cost- effectiveness estimates submitted in response to the appraisal consultation document. The Committee noted that the updated cost-effectiveness estimates incorporated more recent data from the RECORD-1 trial. Therefore it accepted the extension of the time horizon of the model from 144 weeks (18 cycles) to 312 weeks (39 cycles). See FAD sections 3.24 and 4.12

Consultee	Comment	Response
	<ul> <li>been to reduce the ICER. This is because there is greater survival and therefore QALY's in the everolimus arm but no further everolimus treatment costs as these are only applicable for the stable disease states. Full details of these updated analyses follow in the remainder of the document. The results from the PenTAG, RPSFT adjusted analysis and updated Novartis analyses are provided in the following table for ease of comparison.</li> <li>Table – not included here.</li> </ul>	
Novartis	All of the results presented in the above table take into account the patient access scheme (PAS) which was put in place by agreement between Novartis and the DoH, prior to our submission in order to facilitate a positive decision as soon as possible. As this scheme has been approved by the DoH, and is already being implemented by the NHS, the results which incorporate the PAS are the appropriate ones to be considered.	The Committee understood that the cost effectiveness estimates included a patient access scheme which had been agreed with the Department of Health. See FAD Section 4.11
Novartis	The following section summarises an additional body of clinical evidence in order to help the Appraisal Committee decide what constitutes the most plausible estimate of survival in patients receiving BSC following sunitinib failure. The evidence supplied includes the most recent, relevant publications and a survey reflecting UK clinical expert opinion. It should be noted that the reason for conducting the survey was not revealed to the respondents. Finally, because of the lack of directly applicable publications in this area Novartis also requested, and was provided with, primary patient data from clinicians with experience of 1 <sup>st</sup> line sunitinib use to demonstrate what happens to patient's with no 2 <sup>nd</sup> line treatment. Although retrospective in nature, this crucially provides actual UK clinical data from two large London teaching hospitals, The Queen Elizabeth hospital in Birmingham and from two hospital's in the Royal Wolverhampton NHS trust to demonstrate OS in routine clinical practice for patients that received sunitinib therapy and no active 2 <sup>nd</sup> -line therapy following sunitinib failure. A table comparing OS in patients failing on 2 <sup>nd</sup> -line sunitinib is presented below. Table – not included here.	The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the manufacturers' submission, revised cost-effectiveness estimates submitted in response to the appraisal consultation document and corresponding ERG reports. It also carefully considered the comments received from consultees and commentators in response to the Appraisal Consultation Document. The Committee noted that the overall survival associated with best supportive care in the ERG analysis (10.8 months) was likely to be higher

Consultee	Comment	Response
		than in clinical practice. The
		Committee noted that the difference in
		overall survival between treatment
		arms was 8.2 months in the
		manufacturer's revised RPSFT
		analysis and 5.9 months in the ERG's
		revised RPSFT analysis. It noted
		evidence from the manufacturer and
		the clinical specialists that an increase
		in overall survival of 1.4 months per
		month of increased progression-free
		survival would be considered
		plausible. Therefore, the Committee
		agreed that the incremental overall
		survival derived using the
		manufacturer's revised RPSFT
		analyses was almost twice as much
		as would be expected, given that the trial had observed an increase of 3
		months in progression-free survival.
		The Committee accepted that the
		ERG's estimate of the incremental
		difference in overall survival for
		everolimus versus best supportive
		care (5.9 months) was more plausible
		than that derived by the manufacturer
		and it was based on all the available
		data. See FAD section 4.10
Novartis	Because of the lack of prospective clinical data Novartis approached 4 institutions to ask	Comments noted. See detailed
	if they were able to provide data to us for the purpose of verifying in UK clinical practice	response above.
	what the OS was for patients who received 1 <sup>st</sup> line sunitinib and then no further anti-	

Consultee	Comment	Response
	cancer therapy.	
	St Bartholemew's hospital in central London has had gained a lot of experience with sunitinib because of its involvement with the Pfizer expanded access programme and other sunitinib clinical trials. Data from clinical practice which included patients from 2006 to present showed that the median time from CT defined progression to death is 5 months (95% CI 3-7 months). No patient received targeted therapy in the 2 <sup>nd</sup> -line, however some did receive chemotherapy. Patients were excluded from the analyses if they stopped treatment before assessment of disease progression occurred, if they stopped due to toxicity or died on sunitinib or were not assessable for disease progression. It was also noted by the clinician that "most patients continued on sunitinib" even though disease progressed according to RECIST criteria. Novartis feel this is likely to be because of the maintenance of clinical benefit even when disease is progressing according to RECIST criteria.	
Novartis	Novartis would also like to highlight to the Appraisal Committee that patients may have continued to receive sunitinib post-progression due to the lack of alternative active treatment options, especially where the patient maintained performance status, and there was the perception of clinical benefit for the patient and/or clinician beyond RECIST criteria measures.	Comment noted. See detailed response above.
Novartis	The Royal Marsden hospital in South West London also provided retrospective data to Novartis as a result of their long term access to sunitinib through clinical trials and the Pfizer expanded access programs. The data included a total of 87 patients with advanced RCC starting sunitinib at the Royal Marsden Hospital between 2005 and December 2008 that had progressed after an adequate period of treatment and died (not including patients who died on sunitinib treatment or stopped due to toxicity). For the 62 patients who received no 2 <sup>nd</sup> -line therapy the median time to death from stopping sunitinib treatment was 64.5 days (2.12 months). Again considerably shorter than the average 10.9 months OS suggested by the PenTAG model (discounted).	Comment noted. See detailed response above.
Novartis	In addition to this The Queen Elizabeth hospital, Birmingham, provided us with audit data for aRCC patients treated with sunitinib and no further treatment. Due to the centres involvement with sunitinib clinical trials and the expanded access program experience	Comment noted. See detailed response above.

Consultee	Comment	Response
	dated back to 2006 and included patients right up to the present time there was data for 94 patients. For these patient's the median OS was found to be 3.8 months. This data does include patients not yet dead although as there are 23 patients, if these patients are excluded the median overall survival would be much lower.	
Novartis	A clinician with experience of sunitinib use from two hospitals that are part of the Royal Wolverhampton NHS Trust provided data for 8 patients started between 03/2007 and 05/2009. Here the median OS from stopping treatment to death is 2.6 months (2 patients have stopped treatment but remain alive so are not part of this calculation). Patients were not part of the audit if they were taken off treatment due to toxicity or death.	Comment noted. See detailed response above.
Novartis	Finally, Novartis undertook a quantitative on-line survey of clinicians experienced in treating advanced RCC in the UK. No information was provided to respondents about the reason for the survey or who was sponsoring it. Thirty seven clinicians responded to the questionnaire, of these, 26 were consultant grade and 11 were specialist registrars (year 5+) and 34 of the 37 responders were from either teaching hospitals or tertial centres. On average the clinicians treated 33 aRCC patients a year. As Novartis have previously submitted to NICE an estimated eligible patient pool of 982 we believe this covers most of the aRCC population. Novartis feels the sample represents clinicians sufficiently experienced in the treatment of the disease and likely to be involved in prescribing these drugs.	Comment noted. See detailed response above.
	The survey results showed clinicians expected the mean OS after failure on sunitinib with no further active treatment to be 6.1 months (6 months median). 57% of clinicians anticipated the range would be between 6-9 months and only 8% of those surveyed believed OS would be 10-12 months.	
Novartis	There is no published evidence directly in line with the decision problem ie patients who receive BSC only following failure on sunitinib therapy. However, the publication by Di Lorenzo <i>et al.</i> 2009, is informative with regards to OS for 2 <sup>nd</sup> - line patients following sunitinib. The study evaluated the efficacy of sorafenib following failure on sunitinib. The median OS for these patients was 7.4 months. <sup>i</sup> In many respects the patients in this study were reflective of those in the everolimus study (RECORD-1) but patients on the Di Lorenzo study could be considered as having a slightly better prognosis based on the	Comment noted. See detailed response above.

Consultee	Comment	Response
	fact patients in this study generally had better MSKCC profiles which included better performance status and lower rates of metastatic disease in organs such as the liver, lungs and lymph nodes. <sup>i,vi</sup> Considering the fact that these patients were on active anti- tumour therapy and the patients generally had superior prognostic scores, the median OS of 7.4 months might be expected to be a best case scenario or even superior compared to patients who get BSC only following sunitinib. <sup>i</sup>	
Novartis	Finally, Liu <i>et al.</i> presented a poster at European CanCer Organisation/European Society for Medical Oncology (ECCO/ESMO) in September 2009 which retrospectively evaluated patients survival following discontinuation of sunitinib or sorefenib in aRCC patients from routine clinical practice. The median OS results in this study for patients who only received sunitinib was 5.2 months. <sup>ii</sup>	
Novartis	In summary, the preliminary recommendations are based on estimates of cost- effectiveness resulting from PenTAG's adjustments to the Novartis analyses i.e £75,700 and £65,200. However, these estimates are misleading and are unlikely to represent the true value of everolimus. This is because the estimate of £75,700 relies on an estimate of survival in the BSC arm which is unrealistic based on the evidence which has been collated since the Appraisal Committee meeting on the 13 <sup>th</sup> January. In addition, the £65,200 is based on an earlier data cut from the RECORD-1 trial and does not reflect the intended overall, effective mortality HR of 0.55. It is important that the final decision regarding the use of everolimus for aRCC patients should rely on estimates of cost- effectiveness that are based on assumptions of OS in BSC patients that are realistic and consistent with the best available clinical evidence.	Comment noted.
Novartis	We therefore respectfully request that due consideration is given to the additional evidence and revised estimates of cost-effectiveness which take into account all of PenTAG's criticisms, as well as the longer term data available from the RECORD-1 trial. These results demonstrate that everolimus is clinically-effective and based on the end of life criteria, a cost-effective treatment for patients with aRCC who fail on 1 <sup>st</sup> -line sunitinib therapy.	manufacturer's revised cost- effectiveness estimates submitted in

Consultee	Comment	Response
		Therefore it accepted the extension of
		the time horizon of the model from
		144 weeks (18 cycles) to 312 weeks
		(39 cycles). See FAD sections 3.24
		and 4.12
Novartis	Detailed Response to Matters Arising from the Appraisal Consultation Document	Comment noted.
	The preliminary decision not to recommend everolimus is based on the estimates of cost-	
	effectiveness presented by PenTAG. This is because it was felt that Novartis had	
	underestimated OS in the BSC arm using both modelling approaches presented. Based	
	on the information that we have compiled from the published literature, data from routine	
	clinical practice and clinical expert opinion, Novartis strongly believe that the preliminary recommendation is not justified.	
	Therefore based on the above, we do not believe that the provisional recommendations	
	of the Appraisal Committee are sound nor do they constitute a suitable basis for the preparation of guidance to the NHS.	
Novartis	Section A – Main concern	The Committee considered all the
	A1. the preliminary recommendations are based on estimates of cost-effectiveness	evidence submitted, including
	resulting from PenTAG's adjustments to Novartis' analyses i.e £75,700 and	evidence from clinical trials, patient
	£65,200. However, these estimates are misleading and are unlikely to represent the	and clinical experts, the
	true value of everolimus. The reasons for this are as follows:	manufacturers' submission, revised
		cost-effectiveness estimates
	- the estimate of £75,700 from PenTAG's "exploratory analysis" using RPSFT is	submitted in response to the appraisal
Novartis	underpinned by a clinically unrealistic estimate of OS in the BSC arm of 10.9	consultation document and
	months (11.2 months undiscounted). This estimate is therefore unlikely to either	corresponding ERG reports. It also
	represent the most plausible estimate of cost-effectiveness, or reflect the true	carefully considered the comments
	magnitude of survival benefit conferred by everolimus	received from consultees and
		commentators in response to the
		Appraisal Consultation Document.

Consultee	Comment	Response
		The Committee noted that the overall
		survival associated with best
		supportive care in the ERG analysis
		(10.8 months) was likely to be higher
		than in clinical practice. The
		Committee noted that the difference in
		overall survival between treatment
		arms was 8.2 months in the
		manufacturer's revised RPSFT
		analysis and 5.9 months in the ERG's
		revised RPSFT analysis. It noted
		evidence from the manufacturer and
		the clinical specialists that an increase
		in overall survival of 1.4 months per
		month of increased progression-free
		survival would be considered
		plausible. Therefore, the Committee
		agreed that the incremental overall
		survival derived using the
		manufacturer's revised RPSFT
		analyses was almost twice as much
		as would be expected, given that the
		trial had observed an increase of 3
		months in progression-free survival.
		The Committee accepted that the
		ERG's estimate of the incremental
		difference in overall survival for
		everolimus versus best supportive
		care (5.9 months) was more plausible
		than that derived by the manufacturer
		and it was based on all available data.

Consultee	Comment	Response
		See FAD section 4.10
Novartis	- we have been able to show by statistical means that PenTAG's adjustment to the	Comment noted.
	IPCW analysis results in an overall effective HR of 0.6 rather than 0.55. Therefore	
	the estimate of cost-effectiveness of £65,200 is artificially inflated and should not	
	be used as the basis for decision-making. In addition, this estimate is based on the less mature, February 2008 data cut, from the RECORD-1 trial.	
Novartis	B. The current recommendations do not take into account all of the available	The Committee considered all the
	evidence. In addition, the provisional recommendations as detailed in the ACD are	evidence submitted, including
	not justified, nor do they constitute a reliable basis for the provision of sound	evidence from clinical trials, patient
	guidance to the NHS.	and clinical experts, the
		manufacturers' submission, revised
		cost-effectiveness estimates
		submitted in response to the appraisal consultation document and
		corresponding ERG reports. It also carefully considered the comments
		received from consultees and
		commentators in response to the
		Appraisal Consultation Document.
Novartis	B1. The preliminary decision is based on the conclusion that PenTAG's estimates	The Committee noted that the overall
	of cost-effectiveness are more plausible than those presented by Novartis.	survival associated with best
	However, the survival estimate for BSC of 11.2 months (undiscounted) which	
	underpins PenTAG's cost/QALY of around £75,700 is not deemed to be clinically	
	plausible based on the available evidence.	than in clinical practice. The
		Committee noted that the difference in
		overall survival between treatment
		arms was 8.2 months in the
		manufacturer's revised RPSFT
		analysis and 5.9 months in the ERG's

Consultee	Comment	Response
		revised RPSFT analysis. It noted evidence from the manufacturer and the clinical specialists that an increase in overall survival of 1.4 months per month of increased progression-free survival would be considered plausible. Therefore, the Committee agreed that the incremental overall survival derived using the manufacturer's revised RPSFT analyses was almost twice as much as would be expected, given that the trial had observed an increase of 3 months in progression-free survival. The Committee accepted that the ERG's estimate of the incremental difference in overall survival for everolimus versus best supportive care (5.9 months) was more plausible than that derived by the manufacturer and was based on all available data. See FAD section 4.10
Novartis	B1. i) A paper by Di Lorenzo <i>et al.</i> 2009 reports on a study which evaluated the efficacy of sorafenib, in patients who failed sunitinib therapy. <sup>i</sup> These patients were receiving active treatment for their disease as well as being well matched to the RECORD-1 patients with respect to baseline characteristics and, where there were differences, these favoured the sorafenib patients ie the prognostic risk factors such as MSKCC profile, performance status and rates of metastases in liver, lungs and lymph nodes were such that one might expect the patients in the sorafenib study to live longer than those in the RECORD-1 study. This means that survival in the sorafenib patients might be a reasonable and conservative proxy for the BSC patients in the RECORD-1 study. <sup>i,iv</sup> The results from the	Comment noted. See above response

Consultee	Comment	Response
	Di Lorenzo study demonstrated that the median survival in the sorafenib patients was 7.4 months. This is broadly consistent with the estimate of survival from the Novartis RPSFT analysis which estimates a mean survival in BSC patients of 7.9 months, (8.1 months undiscounted). <sup>i</sup>	
Novartis	B1. ii) A poster by Liu <i>et al.</i> presented at the European CanCer Organisation/European Society for Medical Oncology (ECCO/ESMO) in September 2009, evaluated survival rates following discontinuation of sunitinib and sorafenib in aRCC patients in routine clinical practice. <sup>ii</sup> This study involved a retrospective review of data from a US claims database on patients with aRCC. Patients were included in the study if they received sunitinib only, sorafenib only or both treatments and then discontinuation of sunitinib or sorafenib to death. Of the 451 patients identified, 264 patients discontinued treatment and did not restart therapy. Of these 131/264 patients had received sunitinib, 70/264 patients had received sorafenib and 63/264 had received both sunitinib and sorafenib. The median survival in patients who received sunitinib only was 5.2 months. <sup>ii</sup>	Comment noted.
Novartis	B1. iii) As presented in our submission evidence from the literature suggests that, if left untreated, patients with advanced renal cell carcinoma (aRCC) have a limited life expectancy, with a median survival without treatment of 6 to 12 in the first-line setting. <sup>iii,iv,v,</sup> Data from the years prior to VEGF targeted therapy clearly demonstrate that patient's given hormone treatment (medroxyprogesterone) aimed at symptom relief only have a median OS of 6 months. <sup>vii</sup>	
Novartis	There is no direct data published to inform the Appraisal Committee on a patient's OS after TKI therapy if they receive no further active therapy i.e. BSC only 2 <sup>nd</sup> -line. This is mostly because cross over from placebo to active treatment upon progression remained high in trials with targeted agents or because information on PFS is not provided.	Comment noted. See response above.
Novartis	However, patients who are eligible for everolimus will be more advanced with respect to time from diagnosis of aRCC compared to those who have not already failed on at least one previous therapy. <sup>vi</sup> This is important because there is an increasing amount of pre- clinical evidence to suggest that disease may progress more rapidly after resistance	above.

Consultee	Comment	Response
	develops with sunitinib use, <sup>viii</sup> raising the possibility that once patients progress on sunitinib they will have a shorter median OS compared to patients untreated in the 1 <sup>st</sup> line. There is also some limited clinical evidence to support this hypothesis in aRCC clinical practice. In a small, UK clinical study, patients were given chemotherapy after progression on sunitinib. The results of this study for patients with aRCC who had previously been progressed on cytokine therapy and then sunitinib the OS was a median of 4.2 months. <sup>ix</sup>	
Novartis	B1. iv) St Bartholemew's hospital in central London has gained a lot of experience with sunitinib because of its involvement with the Pfizer expanded access programme and other sunitinib clinical trials. Data from clinical practice which included patients from 2006 to current use showed that the median time from CT defined progression to death is 5 months (95% CI 3-7 months). No patient received targeted therapy in the 2 <sup>nd</sup> -line however some did receive chemotherapy. Patients were excluded from the analyses if they stopped treatment due to toxicity or died on sunitinib. It was also noted by the clinician that "most patients continued on sunitinib" even though disease progressed according to RECIST criteria. Novartis feel this is likely to be because of the maintenance of clinical benefit.	Comment noted. See response above.
Novartis	Novartis would also like to highlight a point made in our initial submission that patient's may have continued on sunitinib post-progression due to the lack of alternative active treatment options where the patient remained fit and there was the perception of clinical benefit for the patient and/or clinician.	Comment noted. See response above.
Novartis	The Royal Marsden Hospital in South West London also provided retrospective data to Novartis as a result of their long term access to sunitinib through clinical trials and the Pfizer expanded access programs. The data included a total of 87 patients with advanced RCC starting sunitinib at the Royal Marsden Hospital between 2005 and December 2008 that had progressed after an adequate peroid of treatment and died (not including patients who died on sunitinib treatment or stopped due to toxicity). For the 62 patients who received no 2 <sup>nd</sup> line therapy the median time to death from stopping sunitinib treatment was 64.5 days (2.12 months). Again considerably shorter than the average 10.9 months OS suggested by the PenTAG model (discounted). In addition to this The Queen Elizabeth hospital, Birmingham, provided us with audit data	Comment noted. See response above.

Consultee	Comment	Response
Novartis	for aRCC patients treated with sunitinib and no further treatment. Due to the centres involvement with sunitinib clinical trials and the expanded access program experience	
	dated back to 2006 and included patients right up to the present time there was data for	
	94 patients. For these patient's the median OS was found to be 3.8 months. This data	
	does include patients not yet dead although as there are 23 patients, if these patients are excluded the median OS would be lower.	
Novartis	A clinician with experience of sunitinib use from 2 hospitals that are part of the Royal	-
	Wolverhampton NHS Trust provided data for 8 patients started between 03/2007 and	above.
	05/2009. Here the median OS from stopping treatment to death is 2.6 months (2 patients	
	have stopped treatment but remain alive so are not part of this calculation). Patients were	
Novartis	not part of the audit if they were taken off treatment due to toxicity or death. B1. v) The results from a market research survey demonstrate that 57% of the	Comment noted. See response
Novartis	oncologists surveyed believe, that based on experience, patients live for an average of 6-	above.
	9 months from discontinuation of sunitinib, if left untreated. Compared to only 8% of	
	responders believing that OS would be 10-12.	
Novartis	B1 vi) Section 3.19, page 13 of the ACD referring to the Novartis RPSFT model states,	Comment noted. The Committee
	"The EDC stated that the mentality risk is the best supportive area and been	accepted that the use of a Weibull
	"The ERG stated that the mortality risk in the best supportive care arm had been overestimated." and	distribution, which used all available data, was a more appropriate method
	"The FDC conducted on exploratory analysis using revised transition probabilities for the	for estimating overall survival and
	"The ERG conducted an exploratory analysis using revised transition probabilities for the best supportive care arm of the model."	produced a better fit to the empirical data points. See FAD section 4.9
	The exploratory analysis conducted by PenTAG involved ignoring the last transition probability (cycle 6), calculating a mean of the two previous cycles (cycles 4 and 5) and	
	then applying this value from cycle 6 to cycle 18 in the model. The impact of this revision	
	was to increase the estimated mean survival in the BSC arm from 7.7 months	
	(undiscounted) to 11.2 months (undiscounted). As described above, the latter estimate of	
	mean survival for BSC patients post-sunitinib is likely to be unrealistic whereas the	
	estimate of 7.7 months presented in our original RPSFT model is more consistent with	
	the available evidence base whilst remaining conservative.	

Consultee	Comment	Response
Novartis	In order to address PenTAG's concerns that our estimate of survival is based on a single data point we have presented an updated analysis which uses an average of the last two cycles ie the same approach as that adopted by PenTAG in their exploratory analysis. The results from this analysis give a cost/QALY for everolimus of £49,537. Full details of this updated analysis are provided in Section C1, i) below.	Comment noted. See response above.
Novartis	B1 vii) Section 3.15, page 11 of the ACD referring to the Novartis IPCW model states, "Secondly, the ERG stated that in applying the mortality hazard ratio, the manufacturer overestimated the mortality in the best supportive care arm." In order to correct for this overestimation, PenTAG applied a hazard rate multiplier to each cycle in the model, under the assumption that if each cycle HR equates to 0.55 then the overall effective HR across all cycles would also be 0.55. To evaluate whether the PenTAG adjustment actually results in the intended effect, we analysed the survival curve outputs from the economic model following application of PenTAG's adjustments using regression analysis. A virtual cohort of 20,000 patients (10,000 per arm) were run through the PenTAG adjusted economic model and the resulting survival distributions were analysed using Cox proportional hazards regression with the dependent variable being death and the independent variable being treatment assignment. Patients alive at the end of the model were censored at that point. The results from this analysis demonstrate that the impact of PenTAG's adjustment results in an effective mortality HR of around 0.60 ie mortality in the BSC is underestimated. This was confirmed by an independent health economics statistician from ScHARR, Dr Patrick FitzGerald. Consequently this means that the associated ICER of £65,200 produced by PenTAG is artificially inflated. In order to provide a more robust estimate of cost-effectiveness from the IPCW model, we have updated the analysis to include data from the November 2008 analysis. In addition, we have taken into account PenTAG's criticisms relating to	Comment noted. This error was correct by the ERG in their analysis derived from the revised RPFST analysis (See FAD Section 3.24)

Consultee	Comment	Response
	the transition probabilities and applying the discounting from cycle 2 rather than year 2. In order to check that our approach to applying the 0.55 HR resulted in the desired effect we used the same statistical approach described above and calibrated the results to yield an overall effective HR of 0.55. This confirmed that the implementation of the IPCW method in the model resulted in the intended HR of 0.55. The results from this analysis give a cost/QALY of £52,648. Full details of this updated analysis are provided in Section C1, ii) below.	
Novartis	In summary, as detailed above, the conclusion that PenTAG's estimates of survival in the BSC arm and associated estimates of cost-effectiveness are the most plausible is not supported by the available evidence base nor is it consistent with the views of clinical experts who have experience of treating aRCC. In addition, we have demonstrated through statistical means that the PenTAG adjustment to the IPCW approach did not result in the intended effective mortality hazard ratio of 0.55 but resulted instead in a hazard ratio of 0.60. This means that the PenTAG adjustment underestimated mortality in the BSC arm thus providing an inflated estimate of cost-effectiveness for everolimus. For these reasons, Novartis believe the preliminary decision published in the ACD cannot be considered as sound in the light of the evidence and the draft recommendation does not represent a fair, balanced or evidence based foundation for the provision of guidance to the NHS.	Comment noted
Novartis	C1. Updated Estimates of Cost-effectiveness In order to provide the Appraisal Committee with the most robust and plausible estimates of cost-effectiveness we have updated both of our analyses (RPSFT and IPCW) to take into account PenTAG's criticisms and incorporate the longest term clinical data from RECORD-1 ie data from the November 2008 analysis.	Comment noted. See response below.
Novartis	C1. i) Updated RPSFT Analysis The RPSFT analysis was conducted in response to comments in the Evidence Review Group (ERG) Report that an RPSFT analysis would be preferable to the IPCW approach. Novartis therefore sought and received permission to undertake an RPSFT analysis within a two week timeframe. This initial analysis showed that results from the RPSFT approach were similar to those presented for the IPCW approach. As stated in our submission of the RPSFT results, the limited timeframe did not allow us to add further	The Committee discussed the manufacturer's and ERG's cost- effectiveness estimates derived using the RPSFT method. The Committee considered that the time horizon and discounting in the analyses were appropriate. See FAD sections 4.11

Consultee	Comment	Response
	<ul> <li>cycles to the model to fully capture the additional survival in the RECORD-1 trial demonstrated by the November 2008 data cut as compared to the survival indicated from the earlier February 2008 analysis. In the aforementioned submission we highlighted that the impact of adding further cycles to the RPSFT model would be to decrease the incremental cost-effectiveness ratio due to the additional life years gained (LYG), and therefore QALY's, in the everolimus arm but no further everolimus treatment costs as these are only applicable for the stable disease states. In order to take into account PenTAG's criticisms of our RPSFT analysis and fully incorporate the additional survival benefits demonstrated by the November 2008 data cut the following revisions have been made:</li> <li>the number of cycles in the model have been increased from 18 to 39 in order to capture the greater survival demonstrated by the November 2008 data cut;</li> <li>as suggested by PenTAG, discounting at a rate of 3.5% has been applied from cycle 2;</li> <li>in the BSC arm, for states leading to death, rather than carry the last transition probability forward for the remaining cycles (as in our previous model), we have calculated an average of the transition probabilities from cycles 5 (0.21) and 6 (0.5) and applied this average value (0.35) to all remaining cycles ie cycles 7 to 39.</li> </ul>	and 4.12
	The results from these revisions are presented in the table below.	
	[not reproduced here]	
Novartis	The deterministic results from this analysis give an incremental cost-effectiveness ratio of £49,537 (with PAS). This is underpinned by a mean estimated survival of 7.9 months in the BSC arm and 16.1 months in the everolimus arm. In order to achieve a target threshold of £30,000 per QALY gained, a QALY weight of 1.66 would be required. This QALY weight is within previously accepted limits based on products meeting the end of life criteria.	The Committee discussed the manufacturer's cost-effectiveness estimate of £49,500 per QALY gained which incorporated estimates of clinical effectiveness using the RPSFT method. See FAD Sections 4.10 and 4.11

Consultee	Comment	Response
Novartis	C1. ii) Updated IPCW Analysis	
	The IPCW analysis presented in our original submission was based on data from the	The Committee noted that the key
	February 2008 data cut. The later November 2008 data cut demonstrated greater survival	factor in determining the cost
	in everolimus patients than that indicated by the February 2008 analysis. In order to take	effectiveness was the estimate of
	account of the updated results and take into account PenTAG's criticisms the following	overall survival and discussed the
	revisions have been made to the IPCW analysis:	IPCW and the RPSFT methods used
	- data from the November 2008 data cut has been used to populate the model;	to estimate this from the RECORD-1
	-the number of cycles in the model have been increased from 18 to 39 in order to capture	trial data. It heard from the ERG that it
	the greater survival demonstrated by the November 2008 data cut;	considered the RPSFT method to be
	- as suggested by PenTAG, discounting at a rate of 3.5% has been applied from cycle 2;	more methodologically robust than the
	- as suggested by PenTAG, the HR multiplier has been applied to rates rather than the	IPCW method because it does not
	transition probabilities:	assume that there are no unmeasured
	- as suggested by PenTAG, in the BSC arm, for states leading to death, rather than carry	confounders. In addition, the
	the last transition probability forward for the remaining cycles (as in our previous model),	Committee understood that the
	we have calculated an average of the transition probabilities from cycles 10 (0.24) and 11	manufacturer's revised IPCW analysis
	(0.23) and applied this average value (0.23) to all remaining cycles ie cycles 12 to 39.	contained a number of unexplained
	- overestimation of mortality in BSC arm is corrected by applying the same rate to all	differences between the original and
	transitions leading to death ie from stable disease with adverse events, stable disease	revised models, and so the ERG
	without adverse events and progressed disease to death.	could not conduct a full critique of the
	- transition probabilities were calibrated to ensure an effective HR of 0.55. The effective	revised IPCW analysis. The
	HR was checked by running a virtual cohort of patients through the model and analysing	Committee also noted that the RPSFT
	the survival output using a Cox proportional hazards model.	method had been used previously in
	· · · · · · · · · · · · · · · · · · ·	'Sunitinib for the treatment of
	All other aspects of the model remain unchanged. The associated transition probabilities	gastrointestinal stromal tumours'
	are provided in Appendix 1.	(NICE technology appraisal guidance
		179). The Committee therefore
	The results from the updated analysis are presented in the table below.	concluded that, in this instance, it was
	[not reproduced here]	more appropriate to evaluate the cost
		effectiveness of everolimus based on
		the estimates generated using the
		RPSFT method. See FAD section 4.8

Consultee	Comment	Response
Novartis	The deterministic results from this analysis give an incremental cost-effectiveness ratio of £52,648 (with PAS). This is underpinned by a mean estimated survival of 9.7 months in the BSC arm and 16.2 months in the everolimus arm. In order to achieve a target threshold of £30,000 per QALY gained, a QALY weight of 1.75 would be required. This QALY weight is within previously accepted limits based on products meeting the end of life criteria. In addition, this is likely to be a conservative estimate of cost-effectiveness as it is underpinned by an assumption of survival in the BSC arm which is optimistic based on the available evidence.	The Committee concluded that, in this instance, it was more appropriate to evaluate the cost effectiveness of everolimus based on the manufacturer's revised estimates generated using the RPSFT method. See FAD section 4.8
Novartis	<ul> <li>D. We do not believe that the provisional recommendations as detailed in the ACD are justified nor do they constitute a reliable basis for the provision of sound guidance to the NHS.</li> <li>D1. The decision not to recommend everolimus for the treatment of patients with advanced renal cell carcinoma (aRCC), whose disease has progressed on or after treatment with VEGF-targeted therapy is inappropriate as it relies on the view that the estimates of cost-effectiveness presented by PenTAG are more plausible than those presented by Novartis. This is contrary to the available evidence base.</li> </ul>	Comment noted. See responses above and below.
Novartis	As detailed in Section A1 of this document, Novartis strongly believes the rejection of everolimus for aRCC is perverse in the light of the evidence for the following reasons: - the estimate of survival for BSC patients (11.2 months undiscounted) which underpins PenTAG's estimate of cost-effectiveness is not clinically plausible. This means that the resulting estimate of £75,700/QALY is highly conservative and does not reflect the true value of everolimus; - the estimate of cost-effectiveness of around £65,200/QALY has been shown using statistical means to overestimate survival in the BSC arm thus artificially inflating the cost/QALY.	The Committee noted that the overall survival associated with best supportive care in the ERG analysis (10.8 months) was likely to be higher than in clinical practice. The Committee noted that the difference in overall survival between treatment arms was 8.2 months in the manufacturer's revised RPSFT analysis and 5.9 months in the ERG's revised RPSFT analysis. It noted evidence from the manufacturer and the clinical specialists that an increase

Consultee	Comment	Response
		in overall survival of 1.4 months per month of increased progression-free survival would be considered plausible. Therefore, the Committee agreed that the incremental overall survival derived using the manufacturer's revised RPSFT analyses was almost twice as much as would be expected, given that the trial had observed an increase of 3 months in progression-free survival. The Committee accepted that the ERG's estimate of the incremental difference in overall survival for everolimus versus best supportive care (5.9 months) was more plausible than that derived by the manufacturer and it was based on all the available data. See FAD section 4.10
		The Committee then discussed the manufacturer's and ERG's cost- effectiveness estimates derived using the RPSFT method: £49,500 and £58,300 per QALY gained respectively. The Committee noted its earlier conclusions that the ERG's analysis (which extrapolated the overall survival with best supportive care using all of the available trial data) resulted in the most plausible

Consultee	Comment	Response
		incremental overall survival for everolimus versus best supportive care. The Committee therefore concluded that the ICER of £58,300 per QALY gained (derived by the ERG) was the most plausible. See FAD section 4.11
Novartis	In summary, the preliminary recommendations are perverse in the light of the evidence and accordingly, do not constitute a reasonable or sound basis on which to base guidance to the NHS. In particular, the belief that the estimates of survival for BSC, and therefore cost-effectiveness, are more plausible based on PenTAG's adjustments and exploratory analysis are not supported by the available evidence base or the views of clinical experts. We therefore respectfully request that due consideration is given to the additional evidence and revised estimates of cost-effectiveness which take into account all of PenTAG's criticisms, as well as the longer term data available from the RECORD-1 trial. The revised estimates of cost-effectiveness £49,537/QALY (RPSFT analysis) and £52,648/QALY (IPCW) analysis. These results demonstrate that everolimus is a clinically-effective treatment for aRCC patients with estimates of cost-effectiveness which are within acceptable limits based on previous appraisals for products meeting the end of life criteria.	Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee
Novartis	<ul> <li>E. Other comments</li> <li>Section 3.7, page 7 This section states,</li> <li>"There were more adverse events and serious adverse events (grades 3 to 4) in the everolimus plus best supportive care arm (40.1%) than the placebo plus best supportive care arm (22.6%)."</li> </ul>	Comment noted. The FAD has been amended accordingly. See FAD section 3.8.
	This is confusing as the section refers to both adverse events and serious adverse	

Consultee	Comment	Response
	events. We therefore propose the following amendment,	
	"There were more serious adverse events in the everolimus plus best supportive care arm (40.1%) than the placebo plus best supportive care arm (22.6%)."	
Novartis	Section 3.18, page 13 This section states,	Comment noted. The FAD has been amended accordingly. See FAD section 3.19.
	"This equated to a mean overall survival of 15.18 months with everolimus plus best supportive care and 7.67 months with best supportive care alone (a non-statistically significant gain of 7.51 months)."	
	This statement is misleading as the estimates of survival quoted are generated by the economic model which are not the subject of statistical testing. We therefore propose that the statement is amended as follows,	
	"Estimates of mean overall survival of 15.18 months with everolimus plus best supportive care and 7.67 months with best supportive care alone were generated by the economic model."	
Novartis	Section 4.8, page 17 This section states,	Comment noted. The Committee agreed that the cost estimates used for adverse events in the model were
	"Firstly it did not agree with the assumption that people starting everolimus therapy would all have stable disease without adverse events."	acceptable. See FAD section 4.12
	It is not clear what is meant by this statement. All patients enter the economic model in the stable disease state without adverse events. Once on treatment, some patients will develop adverse events. The rate at which patients move from the stable disease without adverse events health state to the stable disease with adverse events health state is calculated based on patient data from the RECORD-1 study.	

Consultee	Comment	Response	
Novartis	Section 4.13, page 19 This section states,	Comment noted. The Committee discussed the validity of the estimates	
	<ul> <li>"The Committee heard concerns from the ERG that the RPSFT method had been applied incorrectly by the manufacturer. The application of the transition probabilities led to overestimation of the mortality risk."</li> <li>It is not clear why the ERG considered the application of the transition probabilities to be an overestimation of the mortality risk. The RPSFT analysis presented by Novartis resulted in an estimate of mean survival in the BSC arm of around 8 months. Based on the available evidence this estimate of survival is likely to be more plausible than the estimate of mean survival in PenTAG's analysis of around 11 months.</li> </ul>	of overall survival from the manufacturer's and ERG's RPSFT analyses. The Committee noted the ERG's criticism that the manufacturer's extrapolation of long- term survival in the best supportive care arm was still not based on all of the available data (it was based on the mean of cycles 5 and 6 derived from the RPSFT analysis) and that these data may not be representative of the whole trial population. The Committee accepted that the use of a Weibull distribution, which used all available data, was a more appropriate method for estimating overall survival and produced a better fit to the empirical data points. See	
Novartis	Section 4.15, page 20 This section states,	FAD section 4.9 Comment noted. See above response.	
	" The Committee heard that the life expectancy for people with advanced RCC receiving best supportive care alone was unlikely to be greater than 24 months and was potentially as low as 6 months."		
	This statement is misleading. Patients who are eligible for everolimus are those who have failed treatment with sunitinib ie everolimus is indicated for a 2 <sup>nd</sup> line setting. The available evidence suggests that the life expectancy of patients on BSC following		

Consultee	Comment	Response	
	sunitinib failure is likely to be considerably less than 11 months.		

### Comments received from members of the public

Member of the public 1	Appraisal Committee's preliminary recommendations	I strongly disagree with the Appraisal Committees (AC) preliminary recommendations to not recommend Everolimus for the second-line treatment of advanced renal cell carcinoma Kidney Cancer patients have a rare form of cancer and proper consideration has not been given to this important fact. Â This small patient population have access to only one of the ?newer class? of drugs on the market. They are hugely disadvantaged due to the patient numbers being so small. Â Chemotherapy and radiotherapy are not suitable for Kidney Cancer Patients, this leaves access to Sutent as the only possible hope for Kidney cancer patients.	Comment noted. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/TAMetho dsGuideUpdatedJune2008.pdf)
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Member of the public 1	Manufacturer's submission	The actual cost of Everolimus is on a par with Sutent, roughly £100 per day.  Sutent has already been approved by NICE. It is a widely accepted practice in other countries to implement a package of ?Sequential Treatment? for Kidney Cancer patients. If Sutent should stop working a different treatment is used for a period of time and at some point the patient will revert to Sutent. Sequential Treatment is successfully managed in other countries as  a proven and acknowledged method of treating Kidney Cancer. Â If NICE have accepted the cost of treatment with Sutent, this method of ?Sequential Treatment?, which will incur no significant increase in cost, should be an accepted course without further question of cost.	Comment noted. The Appraisal Committee has been given supplementary advice to be taken into account when appraising treatments which may be life extending. This advice addresses the notion of additional benefits not readily captured in the reference case and to have regard to the importance of supporting the development of innovative treatments that are (anticipated to be) licensed for small groups of patients who have an incurable illness. The Committee concluded that everolimus fulfilled the end-of-life criteria and considered it as such.
Member of the public 1		The manufacturers submission and the trials that have taken place demonstrates without doubt that Everolimus is a clinically effective treatment for Kidney Cancer. Â The Evidence Review Group (ERG) have at no point taken into account when commenting on the QALY figures, the fact that the QALY figure was originally introduced nearly 10 years ago and at no point has consideration been made of inflation and increased production costs. At no point is reference made to the ?small? patient numbers involved who will benefit from this treatment, as Kidney Cancer is a rare cancer.	Comment noted. The reference case stipulates that the cost effectiveness of treatments should be expressed in terms of incremental cost per quality-adjusted life year. See Guide to the Methods of Technology Appraisal section 5.2.11 (Available from URL http://www.nice.org.uk/media/B52/A7/TAMetho dsGuideUpdatedJune2008.pdf

Member of the public 1	Consideration of the evidence	?The Committee therefore concluded that although there was sufficient evidence that everolimus increased progression-free and overall survival compared with best supportive care, the magnitude of the overall survival gain was uncertain?.	Comment noted. The Committee noted that the difference in overall survival between patients receiving everolimus and those receiving best supportive care was 8.2 months in the manufacturer's revised RPSFT analysis and 5.9 months in the ERG's revised RPSFT
		It is despicable that because of the small number of patients involved, large clinical trials are impossible in such a short period of time  (whereas in other much larger  patient groups the data available at this stage in a drugs life is somewhat more detailed and far more extensive), this	analysis. It noted the earlier conclusion that an increase in overall survival of 1.4 months per month of increased progression-free survival was plausible. Therefore, the Committee agreed that the incremental overall survival derived using the manufacturer's revised
		very fact is being used against this group as a reason to refuse. Â Large clinical trials are impossible when considering such a small patient group. This is a factor that has clearly not been taken into account, as although  the committee felt that Everolimus is a treatment that is effective, they are refusing on the basis that there is no mass of data available to support this.	RPSFT analysis (8.2 months) was greater than expected, based on the increase in progression-free survival of 3 months observed in the RECORD-1 trial. The Committee accepted that the ERG's estimate of overall survival for patients receiving best supportive care using the RPSFT analysis was higher than observed in clinical practice, but
			the incremental difference in overall survival for everolimus versus best supportive care (5.9 months) was more plausible than that derived by the manufacturer and was based on all the available data. See FAD section 4.10.

Member of the public 1	Implementation	It is accepted that the cost of Everolimus is similar to Sunitinib, it is widely accepted throughout other countries to offer ?Sequential Treatment?, therefore ?additional? costings using this method of Sequential Treatment are not applicable as the cost to the NHS is roughly the same, it is simply a case of switching treatments for a short period.	Comment noted. The Institute recognises that guidance from other organisations may differ from its own guidance, because of different criteria for making decisions. The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the manufacturer's submission and the ERG report.
Member of the public 1	Related NICE guidance	"For over a decade Kidney Cancer patients have had to cope with no new treatments available to beat this aggressive disease. Â After a lengthy battle, Sunitinib was approved by NICE for use by this small patient group. Â But to date no other treatment is available, there is no alternative for patients who cannot tolerate Sunitinib or for patients for whom Sunitinib will not work. Â As Health Secretary Alan Johnson promised this whole area would be addressed and that  NICE would offer greater flexibility for End Of Life Treatments, but NICE have reviewed 5 new drugs and approved only 1 for first line treatment. Additionally there is no evidence that NICE have applied the EOL criteria to this treatment. This small patient group is severely disadvantaged just by having a cancer that is Rare."	Comments noted. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/TAMetho dsGuideUpdatedJune2008.pdf) The Committee took in to account the end of life criteria in reaching its decision. See FAD sections 4.14 and 4.15

Member of the public 1	Proposed date of review of guidance	To consider reviewing the technology in 3 years time when we live in a period of huge dynamic and technological growth in  cancer fighting treatments is archaic, out dated and totally of no use  to this small patient group. The only way to provide the 'magnitude' of results that NICE are requiring is to allow the patients to have access to this treatment. But this is a vicious circle that will never end. Everolimus is to treat a rare cancer, the masses of lengthy trial data will never be gathered due to small patient numbers, and no further data will be allowed to be gathered unless patients are allowed access to this treatment.	Comments noted. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/TAMetho dsGuideUpdatedJune2008.pdf)
Other 1	Appraisal Committee's preliminary recommendations	It is disgraceful, shameful and unethical that NHS cancer patients are denied the opportunity to access treatments recommended by their oncologists and recognised throughout the world and by the UKs medical profession, for reasons of cost, when the NHS spends millions of pounds on treating entirely self-inflicted lifestyle conditions, from obesity and alcoholism to drug addiction and removing tattoos. NHS priorities must change immediately, to allow oncologists to prescribe drugs such as Evarolimus to those patients they consider may benefit in terms of extended lifetime. NICE should immediately approve the use of Evarolimus for patients deemed suitable by their oncologists. Cancer patients should not be condemned to a more premature death than current therapies can prevent, in order merely to allow NHS funds to be diverted to less life-threatening or self-inflicted conditions.	Comment noted. The Institute recognises that guidance from other organisations may differ from its own guidance, because of different criteria for making decisions. The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the manufacturer's submission and the ERG report.

Patient 1	Consideration the evidence	of If Sutent, the only ?End of Life? drug considered acceptable by NICE, either does not work OR cannot be tolerated OR stops working, there is NO alternative treatment. NICE themselves have agreed that chemotherapy OR radiotherapy do not work for kidney cancer.	Comments noted. The Committee agreed that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of
		Everolimus is readily available and routinely used in other EU countries as a second line treatment for mRCC Everolimus is clinically cost effective and is proving to work as we already have patients in the UK successfully taking the drug. Additionally, consider the significant number of patients worldwide who have had access to this treatment for long periods of time and are successfully responding to	everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the
		<ul> <li>the treatment.</li> <li>The cost of Everolimus works out about the same as Sutent 4 weeks active treatment  £100 per day</li> <li>When considering the cost of Everolimus, the small patient numbers who will need it, should be taken into account</li> </ul>	Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/TAMetho dsGuideUpdatedJune2008.pdf)
		more than the actual cost. Our kidney cancer Clinicians want to use Everolimus when Sutent fails. This is referred to as ?sequential treatment? and is a practice successfully used to treat Kidney Cancer patients in other countries. We need the same treatment option here in the UK.	
		Rarer cancer patients are DISCRIMINATED against as their treatments will always be more expensive, due to patient groups being much smaller. This is particularly the case by NICE. Please provide equality, a basic human requirement.	
		I find it absolutely disgusting that NICE with all their significant and costly resources have been considering Everolimus for the past 14 MONTHS. Yet they have only provided Kidney cancer patients just 3 WEEKS to appeal.	47

Patient 2	Appraisal Committee's preliminary recommendations	Complete shock as everolimus works & widely available in EU USA Canada as second line treatment for mRCC. Â Chemo & radiotherapy ineffective for mRCC. Â No other effective alternative treatment for patients after sunitinib or for those unable to tolerate re side effects, high blood pressure etc or when it has ceased to work or doesnt work. Referred to NICE in November 2008 taken too long to assess grossly unfair to give patients only 3 weeks to respond many will be unaware of your decision. Â Typical mRCC patients in their 60s/70s many with no access to computers or not used to Internet. Â Patients totally DISENFRANCHISED from the process regarding life & death decisions made in their name! Â Jargon is that of statisticians engaged at huge cost to taxpayers/stakeholders & to the layman is incomprehensible. Â No level playing field for us to make meaningful comments in appeal. Â System flawed as QALY set at £30,000 max in 1999 no adjustment for inflation since. Â Lack of transparency re your methodology process model & means of calculation of placing value on human life. Up against NHS lobby group CSAS now unfairly influencing appraisal process despite obvious conflict of interest.	Comment noted. The Institute recognises that guidance from other organisations may differ from its own guidance, because of different criteria for making decisions. The Committee considered all the evidence submitted, including evidence from clinical trials, patient and clinical experts, the manufacturer's submission and the ERG report. Comment noted The reference case stipulates that the cost effectiveness of treatments should be expressed in terms of incremental cost per quality-adjusted life year. See Guide to the Methods of Technology Appraisal section 5.2.11 (Available from URL http://www.nice.org.uk/media/B52/A7/TAMetho dsGuideUpdatedJune2008.pdf)
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Patient 2	The technology	It is not clear that the cost models take into consideration generous discounts & free benefits offered by manufacturer. Cost effective analysis model cannot be used for rarer cancer drugs as there is no tested real time comparator available due to the few people with mRCC. Â The overall cost will be lower than sunitinib as there are fewer patients compared with other cancers. Maximum cost approx £4 million pa when NHS wasted £40 million on lawyers fees for abandoned IT project & huge amount wasted on unused drugs.	The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL <u>http://www.nice.org.uk/media/B52/A7/TAMetho</u> <u>dsGuideUpdatedJune2008.pdf</u> )
Patient 2	manufacturer's submission	Everolimus is the only oral mTOR inhibitor thus saving NHS support costs compared to intravenous dosage. Â At 3.3 It is clear that everolimus performs well for those unable to tolerate sunitinib or where it does not work with a 66% reduction in risk of disease progression with everolimus plus best supportive care compared with placebo. Â This in itself justifies NICE recommendation to give ALL patients longer survival prospects. Â At 3.5/3.10 found to DOUBLE overall survival a more significant & valued measure than health related QOL for patients. Â No consideration has been given to Outliers (patients surviving for longer) on everolimus which could skew the Mean in a small patient group. Â The NICE calculation of ICER cannot apply to a small patient group for obvious statistical reasons. Â Calculation of QALY also not consistent as will always be high as there is no netting-off of existing drug as there is none to consider making the process cruel and unfair.	Comment noted. The Committee agreed that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.

Patient 2	consideration of the evidence	The flexible approach promised by last Health Secretary, Alan Johnson, not properly considered. Â We as a small patient group qualify for inclusion in this and we are aware that a cost-effectiveness of up to £70,000 was agreed to be considered in these circumstances. Â However, NICE have increased their estimate from £65,200 to £75,700 per QALY gained in a timely manner therefore lifting us out of this category and denying extra life to many again. Â I cannot believe that these decisions are made with no medical renal oncologist on the Appraisal Committee. Â We will never be able to lift the appalling five-year cancer survival rates in this country to match the EU standard unless patients have access to SEQUENTIAL TREATMENT as commonly seen in many other countries.	The supplementary advice does not suggest for Committee to apply a particular weight for the cost effectiveness estimate to fall within the acceptable threshold range. The Committee is asked to come to a value judgment on whether the magnitude of additional weight, that would need to be assigned to the original QALY benefits in the patient group for the cost effectiveness of the drug to fall within the current threshold range, would be acceptable in light of the evidence presented.
Patient 2	implementation	NICE guidance needs to be made mandatory as already PCTs are adopting differing approaches to sunitinib access	Comment noted.
Patient 2	related NICE guidance	Pazopanib referred to NICE November 2008. Â FAD not expected until December 2010 at earliest. Â As with sunitinib many patients will again die without treatment due to your delay tactics and we pay your salaries!	Comment noted.
Patient 2	proposed date of review of guidance	As with sunitinib many patients will again die without treatment due to your delay tactics and we pay your salaries. We dont seem to be getting through to you people - WE CANNOT WAIT!!	Comment noted.

Carer 1	Appraisal Committee's preliminary recommendations	There is no other effective treatment for patients who cannot tolerate, or who have become immune to Sunitinib effectiveness. There has to be something else for RCC patients to fall back on. Traditional chemotherapy does not work, so patients will be left with no other option other than give up and die. You have already rejected 3 other RCC drugs when Alan Johnson promised greater flexibility from NICE for EOL drugs. I do feel that this is a class decision, patients who can afford to buy privately (like yourselves) will do so, others will not be able to afford it. I feel that that is discrimination!	Comment noted. The Committee agreed that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.
Carer 1	The technology	side effects are much less than Sunitinib and is usually well tolerated.	Comment noted. The Committee was advised by the patient experts and clinical specialists that everolimus would be tolerated by most people with advanced RCC, and the adverse events would not be significantly worse than those experienced with first-line sunitinib therapy. See FAD section 4.3
Carer 1	manufacturer's submission	The evidence is clear that this does work, obviously, if given too late then its overall effect will be less than if given at the right time as decided by a clinician. It is available throughout Europe and other countries and this appraisal goes to show why our survival rates are so low compared with other countries.	Comment noted. The Committee agreed that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.

Carer 1	(consideration of the evidence)	The qualy has not been changed for more than 10 years. To deny a drug that is clinically effective breaches the Human Rights act, it is in effect sentencing patients to an early death. I also note that the committe did not have an RCC specialist there and so specialist knowledge was missing. Surely this is wrong. The input of a specialised clinician is crucial.	Comment noted The reference case stipulates that the cost effectiveness of treatments should be expressed in terms of incremental cost per quality-adjusted life year. See Guide to the Methods of Technology Appraisal section 5.2.11 (Available from URL http://www.nice.org.uk/media/B52/A7/TAMetho dsGuideUpdatedJune2008.pdf)
Carer 1	Related NICE guidance	patients cannot wait for the decision about pazopanib, they need something now.	Comment noted.
Carer 1	(proposed date of review of guidance	what will be different in 2013, if you have the statistics now, why would you change your mind in 2013? Are you just deferring due to cost?	Comment noted. NICE guidance is considered for review typically 3 years after publication of guidance.
Patient 3	Appraisal Committee's preliminary recommendations	I disagree with this decision. Â If Sutent stops working or indeed doesnt work in the first place then Everolimus is the only viable option.	Comment noted. The Committee heard from clinical specialists and patient experts that there are limited treatment options for people with advanced RCC. See FAD sections 4,2 and 4.3.
NHS professional 1	Appraisal Committee's preliminary recommendations	Although the magnitude of the extension of life with this drug are difficult to estimate because of the cross over in the trial these patients are surviving on average 14.78 months. Â This is a significant extension of life which should not be denied to these patients. Â This agent is being used widely across Europe and it is breach of UK patients human rights to deny them access to this medicine.	Comment noted. The Committee discussed the clinical effectiveness of everolimus in people with advanced RCC whose disease had progressed within 6 months of stopping VEGF-targeted treatment. See FAD section 4.4

NHS professional 1	manufacturer's submission	It is important to remember that the median survival of patients treated with interferon as first line therapy for metastatic RCC is around 14 months (without any second line therapy). Â The median survival of patients randomized to second line everolimus was 14.78 months. Â This means that with the use of sunitinib (median survival 26.4 months)followed by everolimus the median survival will be in the order of 41 months i.e. almost three times longer than in the pre-targeted therapy days!	The Committee noted that the difference in overall survival between patients receiving everolimus and those receiving best supportive care was 8.2 months in the manufacturer's revised RPSFT analysis and 5.9 months in the ERG's revised RPSFT analysis. It noted the earlier conclusion that an increase in overall survival of 1.4 months per month of increased progression-free survival was plausible. Therefore, the Committee agreed that the incremental overall survival derived using the manufacturer's revised RPSFT analysis (8.2 months) was greater than expected, based on the increase in progression-free survival of 3 months observed in the RECORD-1 trial. The Committee accepted that the ERG's estimate of overall survival for patients receiving best supportive care using the RPSFT analysis was higher than observed in clinical practice, but the incremental difference in overall survival for everolimus versus best supportive care (5.9 months) was more plausible than that derived by the manufacturer and was based on all the available data. See FAD section 4.10
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Carer 2	Notes	My husband has been on Sutent for over four years and is doing fine. He was told it would work for maybe six months on the RCC he has had since July 2004. Hopefully if he needs it everolimus will do the same but as it is now he will be unable to get it. What right has NICE got to deny anyone the chance to live. They pay for IVF, Cosmetic Surgery and many more expensive not life saving procedures so why deny a relatively small proportion of people this chance.	Comment noted. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL <u>http://www.nice.org.uk/media/B52/A7/TAMetho</u> <u>dsGuideUpdatedJune2008.pdf</u> )
Carer 2	Appraisal Committee's preliminary recommendations	Why when it has been proven to work	Comment noted. The Committee agreed that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.
Carer 2	the technology	If this is not approved drug compays might as well give up on their R & D	Comment noted
Carer 2	(manufacturer's submission)	Statistics never stand up there is always some that do not get the side effects and do well for a greater time	Comment noted
Carer 2	related NICE guidance	Please just pass all these drugs and stop using money on all the unnecessary meetings and paper work which is going on at the moment	Comment noted

Carer 2	(proposed date of review of guidance	The burden that would be lifted off the the already drained patients relatives and carers of this relatively small group of people if this drug could be approved is unimaginable.  Just to give them hope is all that is needed to lift their  spirits and relieve them of the constant worry that is there every minuet of the day and night	Comment noted
Patient 4	Notes	Time to show some backbone and leadership. You are the NI of "Clinical Excellence" The drug works for some people. Now standard treatment in the US and most European Countries, are we always to be the poor mans health service. The decision to prescribe should be down to the Clinicians/Oncologists. It is up to the Government to fund.	Comment noted
Patient 4	Appraisal Committee's preliminary recommendations	Confirms there is no intention of making the NHS cancer services "First Class"	Comment noted
Patient 4	the technology	As it is in tablet form it is cheap to deliver the service to patients.	Comment noted
Patient 4	manufacturer's submission	This proves that it is an effective treatment	Comment noted. The Committee discussed the clinical effectiveness of everolimus in people with advanced RCC whose disease had progressed within 6 months of stopping VEGF-targeted treatment. See FAD section 4.4
Patient 4	consideration of the evidence	A significant number of patients are living with Advanced RCC well over 24 months eg.Myself 5 years	Comment noted.
Patient 4	related NICE guidance	12 months is desperately slow for people with a life threatening disease. 3 months should be the maximum	Comment noted

Patient 4	proposed date of review of guidance	Far too long. Reviews should be every 6 months	Comment noted. NICE guidance is typically considered for review 3 years after guidance is published.
Patient 5	Appraisal Committee's preliminary recommendations	Confirms there is no intention of making the NHS cancer services "First Class"	Comment noted
Patient 5	The technology	As it is in tablet form it is cheap to deliver the service to patients.	Comment noted.
Patient 5	Manufacturer's submission	This proves that it is an effective treatment	Comment noted. The Committee discussed the clinical effectiveness of everolimus in people with advanced RCC whose disease had progressed within 6 months of stopping VEGF-targeted treatment. See FAD section 4.4
Patient 5	Consideration of the evidence	A significant number of patients are living with Advanced RCC well over 24 months eg.Myself 5 years	Comment noted
Patient 5	related NICE guidance	12 months is desperately slow for people with a life threatening disease. 3 months should be the maximum	Comment noted
Patient 5	proposed date of review of guidance	Far too long. Reviews should be every 6 months	Comment noted

Patient 6	Notes	I have rcc stage iv Ive been on sutent now for almost a year, N.I.C.E refused initially to fund sutent but my PCT finally had to agree that my case was justified and allowed funding, shortly before N.I.C.E Â passed it as available to all patients. IT IS WORKING FOR ME, SOME TUMOURS HAVE DIED COMPLETELY AND ALL OTHERS ARE SHRINKING AND BECOMING LESS SOLID SHOWING CYSTIC CHANGES. These drugs work but if sutent should stop working for me I DESERVE THE RIGHT TO BE GIVEN THE NEXT DRUG TO CONTINUE TO BENEFIT FROM THE EXPENSIVE RESEARCH THESE DRUGS HAVE UNDERGONE TO ALLOW US PATIENTS TO LIVE. WE HAVE A RIGHT TO LIFE.	Comment noted. The Committee agreed that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.
Patient 6	Appraisal Committee's preliminary recommendations	IT SHOULD BE ALLOWED, WE CAN ONLY RECEIVE ONE DRUG AT A TIME SO COST IS NOT AN ISSUE.WHAT IS THE POINT OF MILLIONS BEING SPENT ON RESEARCH IF WE ARE NOT PERMITTED TO HAVE THESE DRUGS.	Comment noted
Patient 6	the technology)	COST IS NOT AN ISSUE, OTHER DRUGS COST AS MUCH AND ARE BEING PRESCRIBED.IT WILL ONLY BE GIVEN IF OF BENEFIT SO WE DESERVE THE RIGHT TO BENEFIT FROM THESE NEW DRUGS.	Comment noted
Patient 6	manufacturer's submission	EACH INDIVIDUAL PATIENT WITH GUIDANCE FROM THEIR ONCOLOGIST SHOULD DECIDE ON WHAT IS QUALITY OF LIFE. NO ONE HAS THE RIGHT TO DECIDE ON THE COST OF A LIFE. WE SHOULD ALL DO WHATEVER IT TAKES TO GIVE LIFE WHERE THERE ARE TREATMENTS THAT CAN HELP.	Comment noted
Patient 6	consideration of the evidence	EVIDENCE SHOWS THAT IT CAN EXTEND LIFE, IT SHOULD BE A PATIENTS CHOICE TO HAVE THIS DRUG WITHOUT COST BEING AN ISSUE!	Comment noted

Patient 6	implementation)	IT SHOULD BE PATIENTS WHO GIVE IMPLEMENTATION ADVICE ON DRUGS NOT PEOPLE WHO HAVE VIRTUALLY NO EXPERIENCE OF WHAT WE ARE GOING THROUGH!	Comment noted
Patient 6	related NICE guidance	N.I.C.E GUIDANCE IS OUTDATED AND UNFAIR, WE ARE NOT NUMBERS WE ARE HUMAN BEINGS.THIS GUIDANCE IS BEING PUT INTO ACTION BY DOCTORS AND PROFESSIONALS BUT NOT ONE IS A SPECIALIST IN CANCER, RENAL CELL CANCER OR ONCOLOGY! WOULD YOU LET A BRICK LAYER COOK YOU A 3 COURSE MEAL AND EXPECT A LA CARTE? I THINK NOT!	Comment noted
Patient 6	proposed date of review of guidance	THE PROPOSED DATE FOR REVIEW OF GUIDANCE IS TOO FAR IN THE FUTURE TO BENEFIT PATIENTS NEEDING TREATMENT NOW.	Comment noted
Carer 2	Appraisal Committee's preliminary recommendations	I think once again NICE have let down RCC Patients and left them with no alternative tratments. Â Clearly a life in England is worth nothing. Â If there were alaternative treatments available that were approved by NICE this decision would not be so devastating for patients and their family. Â I cannot understsnd why NICE put finances before approvingany treatment 1st or 2nd line for Kidney cancer patients.	Comment noted
Carer 2	the technology	Actually i do not understand why the medications cost so much and I believe hoe the Drug companies reaches a price needs to be reviewed. Â It is partly this that stop the drug being accessed by patients who need it. Â How is this good business. Â Certainly not for the patients in the UK.	Comment noted
Patient 7	the technology	Why are we putting ££££££ before people. Â Does this drug need to be so expensive ?????	Comment noted

Other 2	Notes	Ok, so NICE admits that Evirolimus/Afinitor does work, but it's too expensive for the NHS to administer, so some people will die early if they can't cough up the dosh. It's the end of the argument for most people who have kidney cancer (maybe?). ///// Q. That's personally fine by me as I'm rich enough to go out and buy these pills if and when I need to, but what about the people who are not as lucky as me? A. They will die earlier than I will, most probably. ///// Q. The NHS was founded on the fundamental principle of free health care for all UK citizens and it strikes me that the top level management of our national health care rationing system cannot broker a better deal with the private industries who do the research, development and marketing of 21st century health care. Are the Government doing a good job given how much taxpayer's money is spent on NICE? A. I don't know, as there is little public evidence either way as far as I can work out, but my MP here in Witney has so far confirmed a casual interest in the Justice for Kidney Cancer Patients campaign. ///// Q. At the moment is NICE promoting a divided society based on wealth? A. As the Churchill nodding dog would say, "Oh Yes". ///// Q. Given that NICE is a public authority and therefore open to scrutiny do they comply promptly to requests under the Freedom of Information Act? A. Absolutely not, and the Information Commissioner will back me up on this issue.	Comment noted. The Committee agreed that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.
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Other 2	Appraisal Committee's preliminary recommendations	<ul> <li>Ok, so NICE admits that Evirolimus/Afinitor does work, but it?s too expensive for the NHS to administer, so some people will die early if they can?t cough up the dosh. It?s the end of the argument for most people who have kidney cancer (maybe?).</li> <li>Q. That?s personally fine by me as I?m rich enough to go out and buy these pills if and when I need to, but what about the people who are not as lucky as me?</li> <li>A. They will die earlier than I will, most probably.</li> <li>Q. The NHS was founded on the fundamental principle of free health care for all UK citizens and it strikes me that the top level management of our national health care rationing system cannot broker a better deal with the private industries who do the research, development and marketing of 21st century health care. Are the Government doing a good job given how much taxpayer?s money is spent on NICE?</li> <li>A. I don?t know, as there is little public evidence either way as far as I can work out, but my MP here in Witney has so far confirmed a casual interest in the Justice for Kidney Cancer Patients campaign.</li> <li>Q. At the moment is NICE promoting a divided society based on wealth?</li> <li>A. As the Churchill nodding dog would say, ?Oh Yes?.</li> </ul>	Comment noted. The Committee agreed that everolimus was likely to be clinically effective. However, for both legal and bioethical reasons the Committee must take account economic considerations and cost effectiveness of everolimus (Social Value Judgments - Principles for the development of NICE guidance; principle 5). The Committee concluded that everolimus would not be a cost-effective uses of NHS resources.
Other 3	implementation	How can you put a price on someones life????????? What it if it was a member of your family??? Would be different then.	Comment noted

Carer 3	Appraisal Committee's preliminary recommendations	There is no second-line treatment option for kidney patients who fail to respond to sunitab and this drug has proven results. Â It should be provided as general treatment and not on the basis of location.	Comment noted. The Committee heard from clinical specialists and patient experts that there are limited treatment options for people with advanced RCC. See FAD sections 4,2 and 4.3.
Member of the Public 2	Appraisal Committee's preliminary recommendations	As only a small number of patients suffer this condition, the drugs are bound to be expensive as the volume is small. By making the recommendation that it is not made available to patients, surely drugs companies are less likely to continue to research cures for this area, as the wont be selling many of these drugs. These drugs may give someone more time with their family and some precious extra life before succumbing to their condition. Please reconsider.	Comment noted. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/TAMetho dsGuideUpdatedJune2008.pdf)
Patient 8	Appraisal Committee's preliminary recommendations	Everolimus has been proven to extend patients lives as a second line treatment and is widely available in other places around the world. As a rarer cancer this treatment will only be appropriate for and used in a relatively few cases and for a relatively short amount of time. Nevertheless this time is invaluable to the patients and their families as, for instance, additional time can be spent with children to prepare them for your death and to make arrangements which will make life easier for everyone. By stating that it is not available it causes untold stress which in itself has a cost attached not just for the patient but also on family and friends. This stress is incurred not just by those who actually need the treatment but also those who may need it some time in the future.	Comment noted. The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/TAMetho dsGuideUpdatedJune2008.pdf)

Patient 8	the technology	If/when sutent fails this has been proven to extend life even if only for a few months (see comments above. The cost is not "until further notice" as it will only work for a limited period of time but that time is invaluable. It is easy to administer and side effects can be controlled.	Comment noted
Patient 8	manufacturer's submission	QALY - the time spent with friends and family preparing one for ones death will help them cope. Please read and listen to the manufacturers submission. There are no other second line treatments for kidney cancer - this is ageist (many kidney cancer patients are over 60 but many are much younger)and sexist (many patients are men although increasing numbers of women are diagnosed now). This would not be condoned for breast cancer. Kidney cancer is seen as an easy target as relatively few people are diagnosed but because of this the sums make sense (rarer cancer). End of life drugs should be available to all.	Comment noted.The Committee's judgements on cost effectiveness are not influenced by the prevalence of a condition. See Guide to the Methods of Technology Appraisal section 6.2.18 (Available from URL http://www.nice.org.uk/media/B52/A7/TAMetho dsGuideUpdatedJune2008.pdf)
Patient 8	consideration of the evidence	Equality issues - breast cancer and other more "attractive" cancers seem to have huge funding poured in whereas if you are a woman suffering with kidney cancer there is relatively little.	Comment noted
Patient 8	implementation	People are dying while this is being debated	Comment noted

Carer 4	Appraisal Committee's preliminary recommendations	3-4 months is the average amount of time, there will and have been patients who it has helped for considerably longer. what are these "limited resources" that NICE refer to every time theres a contentios decision? £1.4 Billion forecast NHS surplus this year enough to fund this? where does it sit with the £40 Million we have spent on legal consultation fees for the non existent IT system? These patients need another option if initial treatment is not tolerated, and pharma companies need to be incentivised to continue to research more efficient drugs that will provide better results in the future. Are we really committed to beating Cancer, especially those rarer cancers? Money and time need to be stripped out of the multi tiered quango structure that comprises PCTs and various advisory boards, and remove the duplication so we can have a faster and more equitable way of ensuring the right drugs are available at the right costs to the Patients that need them.	Comment noted
Carer 4	the technology	I wouls question procurement practises here, and look for some risk sharing initiatives based on other more succesful drugs	Comment noted
Carer 4	manufacturer's submission	The QALY has not been reviewed for over 10 years, how can this still be a relevant benchmark?	Comment noted

Carer 4	implementation	<ul> <li>3-4 months is the average amount of time, there will and have been patients who it has helped for considerably longer. what are these "limited resources" that NICE refer to every time theres a contentios decision? £1.4 Billion forecast NHS surplus this year enough to fund this? where does it sit with the £40 Million we have spent on legal consultation fees for the non existent IT system? These patients need another option if initial treatment is not tolerated, and pharma companies need to be incentivised to continue to research more efficient drugs that will provide better results in the future. Are we really committed to beating Cancer, especially those rarer cancers? Money and time need to be stripped out of the multi tiered quango structure that comprises PCTs and various advisory boards, and remove the duplication so we can have a faster and more equitable way of ensuring the right drugs are available at the right costs to the Patients that need them.</li> <li>3-4 months is the average amount of time, there will and have been patients who it has helped for considerably longer. what are these "limited resources" that NICE refer to every time theres a contentios decision? £1.4 Billion forecast NHS surplus this year enough to fund this? where does it sit with the £40 Million we have spent on legal consultation fees for the non existent IT system? These patients need another option if initial treatment is not</li> </ul>	Comment noted
		forecast NHS surplus this year enough to fund this? where does it sit with the £40 Million we have spent on legal consultation fees for the non existent IT system? These	
		better results in the future. Are we really committed to beating Cancer, especially those rarer cancers? Money and time need to be stripped out of the multi tiered quango structure that comprises PCTs and various advisory boards, and remove the duplication so we can have a faster and more equitable way of ensuring the right drugs are available at the right costs to the Patients that need	
		them.	64

Carer 4	related NICE guidance	Why is this scheduled to take so long? This goes against the 3 month turnaround time for consultation and guidance - please explain - You have had this for over a year already	Comment noted
Carer 4	proposed date of review of guidance	3 week is not long enough to digest the data and return a coherent argument, the initial decision should be made quicker, and patients/carers should have 6 weeks to make a considered response.	Comment noted
Patient 8	Notes	The devastating effect that cancer has on patients and families is boundless. After successful nephrectomies, surely patients are entitled to drugs to benefit them in the future. Kidney cancer is not self-inflicted as other health issues can be, yet they get treated, why cant we? Its disgraceful that drugs are marketed and not available on the NHS to which we have all subscribed over our working years.	Comment noted
Patient 8	Appraisal Committee's preliminary recommendations	Why not? Because of cost, obviously. So why are patients with other health issues being treated and not kidney cancer patients?	Comment noted
Patient 8	the technology	There are side effects to most drugs - live with it if the medication works	Comment noted
Patient 8	manufacturer's submission	Drug companies are making too much money from this - you cannot put a price on ones health and everyone who is eligible, should be given the chance of survival even for only a few more months if that is the case	Comment noted
Patient 8	consideration of the evidence	Any chance for someone to extend their life however long or short, should be offered.	Comment noted
Patient 8	proposed date of review of guidance	2013????? How many will die before then?	Comment noted

### Everolimus for the Treatment of Advanced Renal Cell Carcinoma (aRCC) – Additional Sensitivity Analyses

#### Background

Following the Appraisal Committee meeting on the 11<sup>th</sup> August, a request was received on the 16<sup>th</sup> August to conduct further sensitivity analyses as follows:

- a) Probabilistic Sensitivity Analysis (PSA) to include a plausible range of Overall Survival (OS) estimates;
- b) One way sensitivity analyses to assess the impact of OS on the cost-effectiveness of everolimus;
- c) Further information regarding the estimates of administration time/cost associated with the PAS scheme and inclusion of these estimates in the PSA.

Details regarding these analyses are provided below.

#### a) Probabilistic Sensitivity Analysis (PSA)

A probabilistic sensitivity analysis has been requested by the Appraisal Committee to, "clarify the uncertainty associated with the new evidence on the total costs of use of everolimus in the NHS."

#### **PSA Methods**

Probabilistic sensitivity analysis was performed using a second order Monte Carlo simulation. Two scenarios were explored based on the following ranges for OS:

- Scenario A) OS HR 0.06 to 1.63 representing RPSFT 95% CI (0.5, 8.5) and
- Scenario B) OS HR 0.27 to 0.87 which represents a conservative, relatively more clinically plausible range.

Each analysis comprised 100 iterations using a sample size of 1000 patients per cohort. An overview of the PSA parameters is presented in Table 1.

#### Table 1 - Overview of PSA Parameters

Parameter	Range	Distribution	Justification for Distribution Adopted
Resource costs	0 to infinity	Gamma	Accepted practice for varying costs in the probabilistic sensitivity analysis
PAS administration costs	Mean cost £18.33 (38.29 minutes x £28) plus or minus 1 SE (£4.15)	Gamma	Accepted practice for varying costs in the probabilistic sensitivity analysis
Health state utility values	0 to 1	Beta	Accepted practice for varying utilities in the probabilistic sensitivity analysis
Transition probabilities	Various ranges related to the individual parameter value and sample information	Dirichlet	Accepted practice for varying multinomial data (e.g. more than one possible transition) in the probabilistic sensitivity analysis
RPSFT-derived hazards ratio	Scenario A) OS HR 0.06 to 1.63 ( RPSFT 95% CI (0.5,8.5))	Log Normal	Accepted practice for varying hazards ratio in the probabilistic sensitivity analysis
	Scenario B) OS HR – 0.27 to 0.87*	Log Normal	

\*see below for full description of derivation of this range.

#### Scenario A – RPSFT Confidence Intervals

As acknowledged in Frances Sutcliffe's email dated 16.8.10, the PSA is more complex than usual for the following reasons:

- The 95% confidence intervals are very wide because the RPSFT method preserves randomisation and does not change the level of evidence against the null hypothesis thus resulting in very wide confidence intervals and
- Survival estimates from the RPSFT method have been presented as relative survival rather than the traditional hazard ratios;

As the RPSFT results are expressed as relative survival, hazard ratios corresponding to the 95% CI (0.5, 8.5) were calculated to provide a suitable range for the PSA. In order to do this, Weibull curves were constructed for the BSC arm corresponding to the 95% CI (0.5, 8.5) from the RPSFT analysis. These curves were derived by first calculating transition probabilities using individual patient data from the RPSFT analysis, and then fitting a Weibull distribution to those transition probabilites as described by the ERG for their base-case analysis. The two resulting Weibull curves were used to calculate adjusted transition probabilities which were entered into the base case model to estimate mean life years gained. The modelled estimates of mean OS derived from these curves were then used to calculate hazard ratios (using Excel Solver) of 0.06 and 1.63 (corresponding to relative survival of 8.5 and 0.5 respectively) thus providing a range which could be explored in the PSA. NB: As the hazard ratios relate to the odds of events occurring and the RPSFT results relate to relative survival time the relationship between relative survival and the hazard ratio is not necessarily linear.

#### Scenario B – Analysis to Explore a Relatively More Plausible Clinical Range

Although, the wide 95% confidence intervals produced by the RPSFT analysis are statistically valid, they represent extreme values which are not necessarily clinically plausible. At the lower bound, a relative survival of 0.5 would mean that BSC patients live twice as long as patients on everolimus treatment. This is unlikely to be the case and there is no evidence to suggest that everolimus reduces life expectancy compared to BSC treated patients. The upper bound of 8.5 would mean that patients on everolimus treatment live 8.5 times as long as those on BSC and although this is possible there is no evidence as yet, that this is the case.

We have therefore attempted to define a relatively more clinically plausible range to explore in the PSA. Although there is limited clinical data on the OS of BSC patients, we have defined a range based on the best available data. For the upper limit we have used the point estimate from the November ITT analysis of the pivotal RECORD-1 trial which was described in our original submission. This gives a highly conservative result for the effectiveness of everolimus (HR 0.87) as it includes the 81% of patients randomised to BSC who crossed over to receive active treatment thus confounding the results to the disadvantage of everolimus. In the absence of any other data, it is a reasonable upper bound to use in the PSA for illustrative purposes. In order to define the lower bound we estimated a hazard ratio of 0.27 based on feedback from the clinician survey (described in our ACD response dated, 2.3.10) which suggests that patients on BSC survive on average for 6 months following failure of sunitinib as compared to a median OS of 14.78 months for patients on everolimus from the RECORD-1 trial. Evidence from actual UK clinical audit data (presented in our ACD response dated 2.3.10) suggests that actual OS in BSC patients, post-failure on sunitinib ranges from 2 to 5 months i.e incremental survival would be greater than if BSC patients live for 6 months as assumed in this analysis. A copy of the ACD response is attached as Appendix 1 for your information.

Compared to Scenario A, Scenario B therefore explores a relatively more clinically plausible range in the PSA, albeit conservative, spanning an OS HR of 0.27 to 0.87.

Text or figures underlined and highlighted in yellow are designated as commercial in confidence.

#### **Scenario A PSA Results**

The PSA results relating to Scenario A, are presented in Tables 2 and 3.

### Table 2 - Scatterplot of Costs and Effects for Scenario A, Based on OS HR 95% CI (0.06, 1.63) CIC information removed

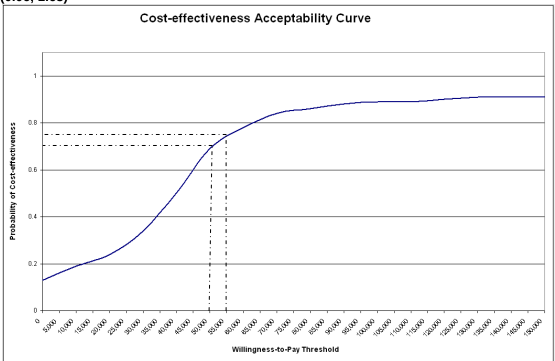


Table 3 – Cost-effectiveness Acceptability Curve for Scenario A, Based on HR 95% CI (0.06, 1.63)

The CEAC demonstrates that there is a 69% probability that the Incremental Costeffectiveness Ratio (ICER) is below £50k and a 73% probability that the ICER is below £55k. However as discussed previously, the wide RPSFT confidence intervals represent extremes which are statistically but not necessarily clinically plausible.

#### **Scenario B PSA Results**

The PSA results relating to Scenario B, are presented in Tables 4 and 5.

## Table 4 - Scatterplot of Costs and Effects for Scenario B Based on OS HR 0.27 to 0.87 CIC information removed

Text or figures underlined and highlighted in yellow are designated as commercial in confidence.

Cost-effectiveness Acceptability Curve

Table 5 – Cost-effectiveness Acceptability Curve for Scenario B) Based on OS HR 0.27 to 0.87)

The CEAC demonstrates that there is a 63% probability that the Incremental Costeffectiveness Ratio (ICER) is below £50k and a 74% probability that the ICER is below £55k. This is likely to be a conservative estimate as the HR defining the upper bound (representing minimum effectiveness of everolimus in terms of effect on OS) is based on the confounded ITT results from the RECORD-1 study where 81% of patients randomised to BSC crossed over to receive everolimus. In addition, the HR defining the lower bound (representing the maximum effectiveness of everolimus in terms of effect on OS) is based on patients on BSC surviving for 6 months, although UK audit data suggests that this may be optimistic. The results demonstrate that the probability of very high ICERs is lower than that suggested by the RPSFT 95% CI.

#### b) One Way Sensitivity Analyses

As requested, one-way sensitivity analyses were conducted around different estimates of OS and PAS administration costs. The results from these analyses are presented in Table 6 below.

Variable	Incremental cost £	Incremental QALY	ICER for everolimus plus BSC versus BSC alone £
Base Case	<mark>xx</mark>	<mark>xx</mark>	49,186
1) Base Case Plus Admin costs (£28)	<mark>xx</mark>	<mark>xx</mark>	49,272

Table 6 – One-way Sensitivity Analysis Results

Text or figures underlined and highlighted in yellow are designated as commercial in confidence.

Variable	Incremental cost £	Incremental QALY	ICER for everolimus plus BSC versus BSC alone £
2) RPSFT 95% CI, 8.5 (HR = 0.06)	24,784	0.734	33,749
3) Assuming OS in BSC arm is 6 months (HR = 0.269)	19,752	0.497	39,724
4) Mean PAS admin cost assuming 2 hours (£ 56)	<u>xx</u>	<u>xx</u>	49,358

Scenario 2) above with an ICER of £33,749 is based on the RPSFT 95% CI, 8.5 which represents everolimus patients living 8.5 times as long as those on BSC. As stated previously as yet there is no evidence that this is likely to be the case and therefore the result from Scenario 2) is likely to be optimistic. Scenario 3) above is based on feedback from a survey of UK clinicians which suggested that on average BSC patients survive for around 6 months. This gives an ICER of £39,724. However, this may be a conservative estimate as data from an audit of UK centres suggested that patients on BSC survived for 2 to 5 months postsunitinib treatment. The results from Scenario 4) demonstrate that even if the mean PAS administration cost was assumed to be 2 hours (mean time from survey = 38 minutes) the ICER would still be below £49,400.

# Resource/Cost Associated with Administering the Everolimus Patient Access Scheme (PAS)

In total ten centres were identified where everolimus patients have been registered using the current PAS. Telephone interviews were conducted with 8 of the pharmacists at centres participating in the PAS. Pharmacists at two of the centres were on vacation and therefore results were obtained from 8/10 of the centres. The summary results from the telephone survey are presented below.

Text or figures underlined and highlighted in yellow are designated as commercial in confidence.

	Time spent on Registration Form in Minutes
Mean (Median)	38.29 (30)
Range	5 to 75
Standard Error	8.89

#### Table of Telephone Survey Results for Time Spent (in minutes) Administering the PAS

The figures in the above table are based on 7/8 of the responses. This was because one of the seven responses was substantially different to the others which were largely consistent. In five out of eight responses the estimated total time to administer the PAS was less than one hour and in two cases around one hour. The remaining response suggested that the scheme would take around eight hours to administer. Importantly, the respondent did comment that this was unlikely to be a robust estimate, due to limited experience of administering the PAS. We therefore consider this response to be an outlier and have consequently not included it in the summary statistics or the PSA. Even if we included the outlier in the summary statistics, the mean time spent administering the PAS would be 94 minutes. Full details of the results from the survey are presented in the appendix attached. Feedback from the survey also suggested that no special qualification, or grade of pharmacist would be necessary to administer the scheme and therefore the costs of administering the PAS are based on an hourly rate of £28 for a basic grade pharmacist (PSSRU 2009). A copy of the draft PAS registration form is attached for your information in Appendix, 2.

#### Conclusions

Results from both PSA scenarios suggest that the probability of everolimus being costeffective at a willingness to pay of £50k/QALY is at least 63% and at a willingness to pay of £55k/QALY is at least 73%. Furthermore the results from Scenario B, which represents a relatively more clinically plausible range, demonstrates that the probability of very high ICERs is lower than that suggested by the RPSFT 95% CI. As Scenario B, is based on conservative estimates of the effectiveness of everolimus, particularly with respect to the HR of 0.87 at the upper bound, it is likely that the true cost-effectiveness of everolimus is likely to be better than that suggested by the PSA in either scenario.

Respondent	At point of registration	Within 2 months of registration	Beyond 2 months of registration	Total time (mins)
<mark>1</mark>	x 🔒	x	×	×
<mark>2</mark>	x 🖌	x	×	×
<mark>3</mark>	x 🖌	x	<mark>x</mark>	×
<mark>4</mark>	x 🛔	x	×	×
<mark>5</mark>	x	x x	× ×	× ×
<mark>6</mark>	x 🛔	x	×	×
<mark>7</mark>	x	x	×	×
<mark>8*</mark>	x 🔒	x	x	
			total time (mins)	275.0
			mean	39.29
			median	30.00
			standard deviation	23.53
			standard error	8.89
			min time	15
			max time	75

#### Appendix – Results from Survey to Ascertain PAS Administration Time

\* not used in the analysis

#### Dear Lori

As requested in the email below, the mean probabilistic ICERs are as follows: Scenario A) RPSFT range based on CI 95% (OS HR 0.06 to 1.63) - mean probabilistic ICER = £50,047 Scenario B) based on OS HR 0.27 to 0,87 - mean probabilistic ICER = £47,811 As discussed between and Rebecca Trowman the 95% confidence intervals are no longer required. A copy of the model is attached for your information. Please treat this model as commercial in confidence. If you have any questions or require any further information please do not hesitate to contact me.

Kind regards

#### <u>Everolimus for the Treatment of Advanced Renal Cell Carcinoma (aRCC) –</u> <u>Details of Updated Patient Access Scheme (PAS) and Associated Results</u>

#### Background

The decision not to recommend everolimus for advanced renal cell carcinoma, as detailed in the Final Appraisal Determination (FAD) dated June 2010, is based on a cost/QALY of £58.3k. This estimate of cost-effectiveness was generated from the ERG's adjustments to the Novartis model and the incorporation of a PAS which was approved by the Department of Health prior to the first Appraisal Committee meeting. In order to expedite the availability of everolimus to patients with advanced renal cell carcinoma, rather than submitting an appeal, we have decided to offer a revised PAS which will further reduce the cost of everolimus to the NHS. The original PAS provides the first pack free with a 5% discount on list price applied to all subsequent packs.

#### The Updated PAS

#### Original PAS

 $1^{st}$  pack (10mg or 5mg tablets x 30) at zero cost to the NHS. Subsequent packs (10mg x 30) will be offered to the NHS at a cost of £2,822.

The list price is £2,970 per pack of 30 x 10mg tablets.

Updated Costs Applied in the Mode	I (without dose intensity adjustment)

Everolimus Cost with Original PAS			Everolimus Cost with Updated PAS			
Unit cost (30 x 10mg tablet pack) £	Total cost per 8 week cycle £	Total cost per 8 week cycle– subseque nt cycles £	Unit cost (30 x 10mg tablet pack) £	Total cost per 8 week cycle– first cycle £	Total cost per 8 week cycle– second cycle £	Total cost per 8 week cycle– subsequent cycles £
2822.00	2445.30	5266.80	xxxxx	<mark>x</mark>	xxxxx*	xxxxxx

The dose intensity adjustment of 91.8% is applied automatically within the model and therefore the associated figures are not supplied in this document.

#### **Summary of Results**

The results of incorporating the updated PAS are presented in the following table.

	Everolimus plus BSC QALY	BSC alone QALY	Inc QALY	Everolimus plus BSC cost (£)	BSC alone cost (£)	Inc cost (£)	ICER for everolimus plus BSC versus BSC alone (£/QALY)
ERG Model with Original PAS	0.843	0.517	0.33	33,854	14,868	18,986	£58,316
ERG Model with Updated PAS	0.843	0.517	0.33	xxxx	xxxxx	xxxxxx	£49,186
ERG Model with Updated PAS and admin cost*	.843	0.517	0.33	xxxxx	xxxxx	xxxxxx	£49,272

#### Table of Cost-effectiveness Results

\*An assumed administration cost has been added to the model allowing for one hour of pharmacist time.

#### Description of Changes to the Model to Take Account of the Updated PAS

The transition probabilities from the ERG's model and supplied by Rebecca Trowman on 1.7.10 were applied in the Novartis model. This enabled us to reproduce the £58,316 estimate. This amended model was then used to calculate the impact of the updated PAS on the cost/QALY.

- In the "Deterministic Results" worksheet the "Patient access scheme: 30 days Free" box was unchecked. NB: Checking this box removes the cost of 30 days treatment only, thus leaving 26 days worth of Afinitor treatment costs in cycle 1. Therefore it is important that this box is unchecked to ensure that a full 56 days worth of Afinitor treatment costs are removed from cycle 1.
- 2) In the "Markov Model (Afinitor)" worksheet, the formula in cell Q8 (costs, undiscounted, cycle 1) was changed from,

=IF('Deterministic Results'!\$W\$13=TRUE,Q7+C8\*((cAfinitorTx/2.154)+cSD)+G8\*((cAfinitorTx/2.154))+cSD+cAfinitorAE)+J8\*\$D\$51+(L8-

 $\label{eq:linear} L7)*D$52,Q7+C8*((cAfinitorTx)+cSD)+G8*((cAfinitorTx)+cSD+cAfinitorAE)+J8*\\ $D$51+(L8-L7)*D$52)$ 

to

=IF('Deterministic Results'!\$W\$13=TRUE,Q7+C8\*((cAfinitorTx/2.154)+cSD)+G8\*((cAfinitorTx/2.154)+cSD+cAfinitorAE)+J8\*\$D\$51+(L8-L7)\*\$D\$52,Q7+(C8\*cSD)+G8\*(cSD+cAfinitorAE)+J8\*\$D\$51+(L8-L7)\*\$D\$52)

This removes all Afinitor costs from cycle 1.

3) In the "Markov Model (Afinitor)" worksheet, the formula in cell Q9 (costs, undiscounted, cycle 2) was changed from,

```
=Q8+C9*$K$49+C8*D9*$D$50+G8*(1-
SUM(H9:I9))*$K$50+(C8*E9+G8*H9)*$D$51+J8*(1-K9)*$K$51+(L9-
L8)*$D$52
```

to

=Q8+C9\*((cAfinitorTx/1.076)+cSD)+C8\*D9\*((cAfinitorTx/1.076)+cSD+cAfinito rAE)+G8\*(1-SUM(H9:I9))\*((cAfinitorTx/1.076)+cSD)+ (C8\*E9+G8\*H9)\*\$D\$51+J8\*(1-K9)\*\$K\$51+(L9-L8)\*\$D\$52

No other changes were made to the ERG adjusted model. Implementation of the above changes results in a revised cost/QALY of £49,186.

The addition of a further  $\pounds 28$  in cycle 1, to cover the cost of 1 hour of pharmacist time was added as an assumed cost of administering the scheme. This increased the ICER from  $\pounds 49,186$  to  $\pounds 49,272$ .

#### Impact of Updated PAS on NHS Administration Requirements

As this is a finance based scheme, there is no requirement to track individual patients. In addition, we believe that the proposed changes to the PAS have negligible additional impact on the level of administration required. This is because we will be issuing the first two packs of Afinitor together rather than just the first pack. The discount on subsequent packs will be applied at the point of invoice. No guidance has yet been provided by the DoH or PASLU as to the likely NHS administration requirements but in view of the simplicity of the scheme an assumption of 1 hour of a pharmacist's time seems reasonable. The impact of adding

the cost of 1 hour of a pharmacist's time into the model is to increase the ICER from  $\pounds$ 49.1k to  $\pounds$ 49.2k.

#### Summary response of PenTAG to Novartis submission:

# "Everolimus for the Treatment of Advanced Renal Cell Carcinoma (aRCC) – Updated Patient Access Scheme (PAS) and Associated Results"

#### Background

In response to the recent Final Appraisal Determination by NICE (dated June 2010) not to recommend everolimus for advanced renal cell carcinoma, Novartis have recently proposed a revised Patient Access Scheme (PAS). The effect of this updated PAS on the cost-effectiveness analysis is examined in more detail below.

#### The Updated PAS

It is clear that the updated PAS proposed by Novartis has the effect of reducing the overall cost of everolimus treatment (in comparison with the original base case cost of the original PAS). The effect of the updated PAS therefore is to reduce the overall ICER output from the cost-effectiveness model. The relationship between the cost of everolimus and ICER as output by the original model is shown in Figure 1 below.

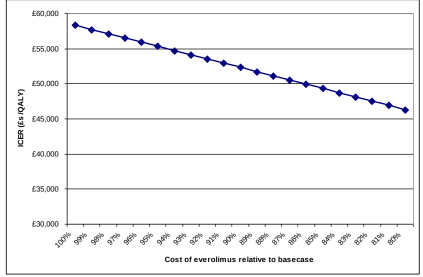


Figure 1: Effect of everolimus treatment cost on model ICER

(ICER shown with original PAS applied. Costs and benefits discounted at 3.5%)

#### Results

We examined the changes to the model as outlined in the Novartis document and were able to replicated their deterministic results using our model (as shown in Table 1 below).

We were also satisfied that the changes to the cost-effectiveness model as outlined in the Novartis submission appropriately represented the conditions of the updated PAS.

	Everolimus plus BSC QALY	BSC alone QALY	Inc QALY	Everolimus plus BSC cost (£)	BSC alone cost (£)	Inc cost (£)	ICER for everolimus plus BSC versus BSC alone (£/QALY)
ERG Model with Original PAS	0.843	0.517	0.33	33,854	14,868	18,986	£58,316
ERG Model with Updated PAS	0.843	0.517	0.33	xxx	xxx	xxx	£49,186
ERG Model with Updated PAS and admin cost*	.843	0.517	0.33	xxx	<mark>xxx</mark>	xxx	£49,272

 Table 1: Cost-effectiveness Results (after Novartis submission)

\*An assumed administration cost has been added to the model allowing for one hour of pharmacist time.

#### Comments

Having examined the Novartis submission we are confident that the adjustments made to the model are appropriate given the proposals outlined in the updated PAS and that these changes have been properly implemented in the model to produce the deterministic outputs as outlined in Table 1 above.

We would however caution that no sensitivity analyses (either one-way or probabilistic) have been supplied with this recent submission. This is against a background of considerable uncertainties known to be inherent in the data used in the model, particularly in relation to the methods used derive the hazard ratio for overall survival between the model arms.

As stated in previous PenTAG submissions, therefore, we would consider a comprehensive probabilistic and one-way sensitivity analysis of model outputs to be essential to inform policy making in this area.

## Everolimus for Second-line Treatment of Advanced Renal Cell Carcinoma (aRCC)

#### ERG response to Additional Sensitivity Analyses.

#### Introduction

In response to requests from the NICE Appraisal Committee on 11<sup>th</sup> August, Novartis have presented additional sensitivity analyses. These include a Probabilistic Sensitivity Analysis (PSA), one-way sensitivity analyses, and further information relating to administration of the updated Patient Access Scheme.

The documents outlining these analyses were received by PenTAG (the ERG) on 21 Sept 2010 and a copy of the model used for the PSA was subsequently received on 28 Sept 2010. Given the clear time constraints entailed it was not possible to perform a full and thorough evaluation of the model and presented analysis before preparing this document. A summary evaluation of the Novartis submission however is given below.

#### Analysis of Probablistic Sensitivity Analysis

In the PSA analysis Novartis present two alternative scenarios based on the following two confidence intervals (CIs) for the hazard ratio for overall survival:

- the CIs derived from the RPSFT analysis on which the overall survival curves are based (HR 95% CI : 0.06, 1.63).
- CIs for the hazard ratio deemed more 'clinical plausible' by Novartis (HR 95% CI: 0.27, 0.87)

We examined both these sets of analysis. We were surprise to see that no summary statistics relating to the mean incremental costs and benefits in the PSA had been provided in the Novartis presentation (these were subsequently reported by email). We were also surprised to see that relatively few iterations (i.e. 100 trials) of the simulation had been used in the PSA. We therefore re-ran the model with 1000 iterations to test the model and results provided by Novartis. The results from these re-runs are presented below and are less favourable to everolimus than the results reported in the Novartis submission.

During our examination of the model we noticed a clear error in the CEAC presented for Scenario A in the Novartis submission (Table 3 in the Novartis submission). This shows a positive probability that everolimus is cost-effective at a zero willingness-to-pay threshold which would imply that in some simulation trials, treatment with everolimus plus BSC costs less than BSC alone. On examination we found that this error is cause by the fact that the Excel model sheet includes dominated trial outputs in the total of ICERs at zero (Cell: "Per Patient Model Results"!BM8). This entails in turn that they contributed to the probability of the ICER falling below £50K/QALY which falsely increases the reported probability of everolimus falling below the £50K/QALY willingness to pay threshold in Scenario A. Our results below correct for this error.

A number of other observations made from our examination of the model and documentation are listed here:

- We are not clear that all sources of potential uncertainty had been included in the model. For example we could find no evidence that the uncertainty surrounding the Weibull survival curve fit had been incorporated.
- The choice of distributions seems broadly in-line with accepted practice, however, the key parameters to determine the shape of probability distributions are not properly reported. The range information given in Table 1 is insufficient (for instance all Beta

distributions have a range of 0-1). These key parameters should ideally be specified as well as their source and justification.

• The method of implementing the PSA by applying probabilities to individual patients (i.e. a per patient approach) is unusual and we did not fully understand why the more conventional cohort based approach had not been adopted.

#### PSA Results from Scenario A (re-run with 1000 simulation trials)

Based on Overall Survival confidence limits derived from RPSFT analysis

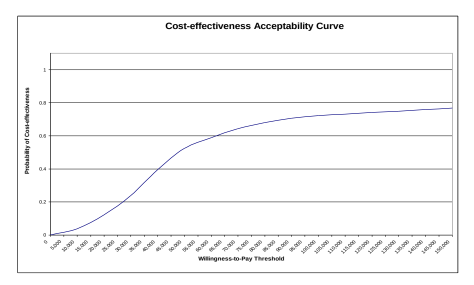
Table 1 : Scenario A - summary PSA outputs

Mean inc.	Mean inc.	ICER	Mean Net	Prob.	Prob. ICER
Costs	QALY	£s/QALY	benefit @	ICER <	< £50K
			£50K/QALY	£30K	
XXX	XXX	£51,661	- £440.77	24.0%	52.7%

Figure 1 : Cost Effectiveness Scatterplot for Scenario A (CIs based on RPSFT HR 95% 0.06, 1.63)

CIC removed

Figure 2 : Cost-effectiveness Acceptibility Curve for Scenario A (CIs based on RPSFT HR 95% 0.06, 1.63)



#### PSA Results from Scenario B (re-run with 1000 simulation trials)

Based on Overall Survival confidence limits suggested by Novartis

Table 2 : Scenario B - summary PSA outputs

Mean Inc.	Mean inc	ICER	Mean Net	Prob.	Prob. ICER
Costs	QALY	£s/QALY	benefit @	ICER <	< £50K
			£50K/QALY	£30K	
XXX	XXX	£49,479	£152.72	28.0%	52.6%

### Figure 3 : Cost Effectiveness Scatterplot for Scenario B (CIs suggested by Novartis OS HR 95% 0.27, 0.87)

CIC removed

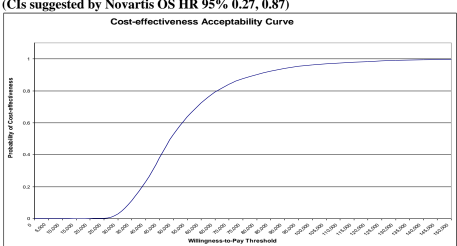


Figure 4 : Cost-effectiveness acceptibility curve for Scenario B (CIs suggested by Novartis OS HR 95% 0.27, 0.87)

#### One-way sensitivity analysis

In our view the one-way sensitivity analysis provided by the Novartis submission is clearly inadequate. Four separate analyses are presented but these look only at the impact of changes in one direction from the base case. Normal practice is to examine the impact of changes to each parameter of interest in both directions and typically to examine a number of different levels for each parameter value. Despite this these one-way analyses confirm once again the centrality of the overall survival hazard ratio in driving the model ICER.

#### Conclusion

The PSA presented by Novartis confirms the high levels of uncertainty associated with the base case estimate of cost-effectiveness for everolimus and BSC vs BSC alone for aRCC. The primary driver for uncertainty in the model outputs is the uncertainty surrounding the estimate of hazard ratio for overall survival between arms.

Two separate analyses are presented by Novartis based on different assessments of the confidence limits that should be applied to the hazard ratio for overall survival. These give slightly different levels of probability that the treatment is cost-effective at the £50,000 per QALY willingness to pay threshold as well as slightly variant mean levels for the incremental costs and benefits.

When we re-ran the Novartis model with a 1000 simulation trials and we needed to correct for an error in the CEAC calculation. We found that mean levels from the PSA analysis of incremental cost and benefit gave an ICER very close to the threshold level of £50,000 per QALY. The CEACs also show that the probability that treatment with everolimus versus BSC is cost-effective at this threshold is very close to 50% in both scenarios.

Constraints in the time available to assess the Novartis PSA analysis have meant that a thorough and in-depth evaluation has not been possible however we would point to the fact that our overview found at least one basic error (i.e. the calculation of the CEAC for Scenario

A) and some clear reporting omissions in the Novartis submission (e.g. too few simulation trials, lack of PSA means, lack of detail, and lack of comprehensive one-way sensitivity outputs). These failings raise questions about the general confidence that can be given to the overall findings presented by Novartis.