### National Institute for Health and Clinical Excellence Centre for Health Technology Evaluation

#### **Pro-forma Response**

#### **Executable Model**

# Bortezomib and Thalidomide for the first-line treatment of multiple myeloma

The economic model enclosed and its contents are confidential and are protected by intellectual property rights, which are owned by **Southampton Health Technology Assessment Centre**. It has been sent to you for information only. It cannot be used for any other purpose than to inform your understanding of the appraisal. Accordingly, neither the model nor its contents should be divulged to anyone other than those individuals within your organisation who need to see to them to enable you to prepare your response. Those to whom you do show the documents must be advised they are bound by the terms of the Confidentiality Acknowledgement and Undertaking Form that has already been signed and returned to the Institute by your organisation.

You may not make copies of the file and you must delete the file from your records when the appraisal process, and any possible appeal, are complete. You must confirm to us in writing that you have done so. You may not publish it in whole or part, or use it to inform the development of other economic models.

## The model must not be re-run for purposes other that the testing of its reliability.

Please set out your comments on reliability in writing providing separate justification, with supporting information, for each specific comment made. Where you have made an alteration to the model details of how this alteration was implemented in the model (e.g. in terms of programme code) must be given in sufficient detail to enable your changes to be replicated from the information provided. Please use the attached pro-forma to present your response.

Please prepare your response carefully. Responses which contain errors or are internally inconsistent (for example where we are unable to replicate the results claimed by implementing the changes said to have been made to the model) will be rejected without further consideration.

Results from amended versions of the model will only be accepted if their purpose is to test robustness and reliability of the economic model. Results calculated purely for the purpose of using alternative inputs will not be accepted.

No electronic versions of the economic model will be accepted with your response.

Responses should be provided in tabular format as suggested below (please add further tables if necessary).

March 2011

#### Issue 1 Cost of managing adverse events

Description of problem	Description of proposed amendment				Result of amended model or expected impact on the result (if applicable)				
The cost of managing adverse events (AEs) was retrieved from TA171. The following	To be consistent with TA171, the following values were used to populate the cells Y30:AD30 on the Costs sheet:						The model was corrected for all the inconsistencies identified – the correspondi		
inconsistencies have been found:	Grade 3 Grade 4					1	results after adjustments are reported below		
<ul> <li>Distribution of management settings of grade 3/4 neutropenia (Y30:AD30, Costs sheet) was not the same as in the quoted source (p. 69 of the ERG report of TA171)</li> <li>In TA171, AEs could be managed in primary care or community care settings, which was not accounted for in the SHTAC model., Although the cost of treating AEs was under-estimated in the model, this was not acknowledged in the report.</li> </ul>	Neutropenia IP: inpatient; DC: day of As the data were to infection from TA1 values. Due to a lack of da community care w	Grade 3Grade 4IPDCOPIPDCOPNeutropenia5.00%55.56%39.44%12.80%42.00%45.20%IP: inpatient; DC: day case; OP: outpatientAs the data were not available for dizziness/fatigue and infection from TA171, no change was made to these values.Due to a lack of data, the cost of primary care and community care was not included in the model.					In issue 4.		

#### Issue 2 Treatment duration

Description of problem	Description of proposed amendment	Result of amended model or expected impact on the result (if applicable)			
The long term efficacy data for MPT were taken from the trials reported by Facon and Hulin. For consistency, the treatment duration used in the model should be set to the mean duration observed in these two trials. Similarly, as the VMP efficacy data are based on the VISTA trial, the treatment duration observed in VISTA should be used in the model.	In the trials reported by Facon and Hulin, patients on MPT had a median treatment duration of 11 and 13.5 months respectively. The number of MPT cycles was therefore set to 10 cycles to reflect a mean treatment duration of 53 weeks. Based on the number of vials actually used in the VISTA trial, the average number of vials was set to 31.5. In the model, the treatment duration was set to 4 cycles for VMP to reflect the average of 31.5 vials, as this was done in the SHTACs updated analyses incorporating this change.	The model was corrected for all the inconsistencies identified – the corresponding results after adjustments are reported below in issue 4.			

#### Issue 3 Estimation of QALYs

Description of problem	Description of proposed amendment	Result of amended model or expected impact on the result (if applicable)			
In the SHTAC model, QALYs were estimated by multiplying the duration of staying in a state by the corresponding health-state utility value (HSUV) instead of using the Markov trace	The Markov trace was generated for pre-progression on treatment (PFS), pre-progression off-treatment (PFS – pre-progression on treatment), progressed (OS – PFS) and died (1 – OS). The probability of being in each state was then multiplied by the HSUV, and then multiplied to cycle length in order to estimate the number of QALYs.	The model was corrected for all the inconsistencies identified – the corresponding results after adjustments are reported below in issue 4.			

#### Issue 4 Cost of the second-line treatment

Description of problem	Description of proposed amendment	Result of amended model or expected impact on the result (if applicable)			
<ul> <li>The following inconsistencies have been found:</li> <li>The SHTAC model assumed a basket cost for the second-line treatment and all patients who progressed were assumed to incur such cost. This approach does not account for the duration of the second-line treatment and results in an overestimation of the cost when a discount rate is applied.</li> <li>A half-cycle adjustment was applied to the cost of the second-line treatment – the model applied an increment of 0.5 to the discount rate (cell K40 on VMP sheet and to cell K39 on other results sheets), while the increment should be of half a cycle (0.5*42/365=0.058 year).</li> </ul>	The cost of the second-line treatment was re-estimated based on the protocol treatment duration, proportion of patients who received the second-line treatment and the split between treatments. For the half-cycle adjustment, 0.5*42/365 was used instead of 0.5	After adjustment for these inconsistencies, the ICER of MPT vs. MP increased from £9,174 to £11,511 per QALY gained.			

#### Issue 5 Selection of evidence

Description of problem	Description of proposed am	Result of amended model or expected impact on the result (if applicable)					
Despite the claim in the SHTAC report of the economic analysis that where possible values from the pre-maintenance phase of maintenance studies was used, this is clearly not the case in the model. The cells of the model F6 and potentially F7 on the 'trial data' sheet should have included a value based on the full trial data, as these relate to periods unaffected by maintenance. Had this been done it would not have been necessary for the committee to take these maintenance studies into account 'but not give them undue weight' as they would have been weighted only for the period of induction MPT treatment in the economic analysis. Given the stated equivalence of MPT and CTDa the same approach should have been followed for this comparison with the MMIX data used for the first induction period and MPT data thereafter.	The hazard ratios were re-estima meta-analysis for the first year as	After applying these data to the updated model, the results were as follows:					
	<ul> <li>The studies by Hulin and F over the full follow-up period</li> </ul>	Scenario	MPT vs. MP	VMP vs. MP	CTDa vs. MP	VMP vs. MPT	
	<ul> <li>over the full follow-up period</li> <li>The "maintenance studies" Gulbrandsen) were consider duration of the induction phy weeks respectively. As this two cycles in the model, the according to two scenarios the estimates from the main the first 6-month period and using the estimates for the</li> <li>For CTDa, the hazard ration were applied for one or two the selected scenario, and observed in the MPT arm wo The previously described a treatment for MPT was also The following hazard ratios wo OS vs. MP</li> </ul>	Corrected* Corrected & adjusted for 1 cycle** Corrected & adjusted for 2 cycles** Dom'ed: domi * Issues 1 to 4 ** Issues 1 to	MP f11,511 f12,893 f13,722 nated accounted 5 accounted	MP           £19,505           £19,505           £19,505           Ifor           d for	MP £34,014 £11,890 £14'677	MPT £211,508 £46,572 £36,794	
	6 - 120.73OS: overall survival; PFS: progression-free						