

Modelling Overall Survival in Immunotherapy Using Parametric Techniques: Avelumab in Previously Treated Metastatic Merkel Cell Carcinoma

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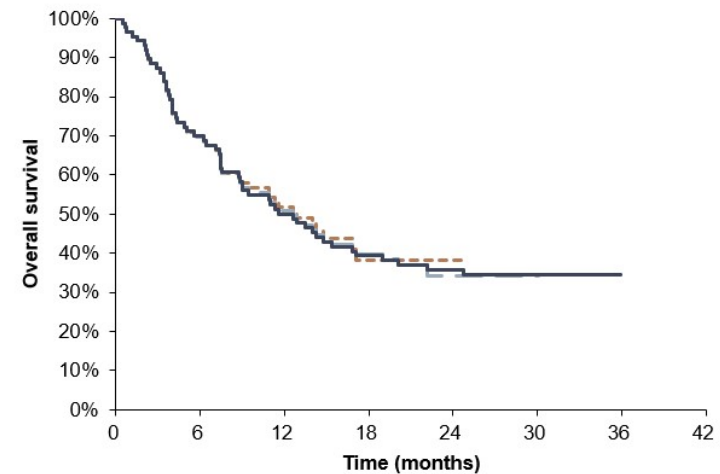
Background and Objectives

- Avelumab (an anti-PD-L1 immune-checkpoint inhibitor) was recently approved in the United States, Europe, and Japan, among others, for the treatment of patients with metastatic Merkel cell carcinoma (mMCC)
- Safety and efficacy data are available from the JAVELIN Merkel 200: Part A trial of 88 patients with previously treated mMCC (NCT02155647)
- The availability of increasingly maturing data from JAVELIN Merkel 200: Part A allows for the production (and subsequent validation) of overall survival (OS) extrapolations
- This analysis compares observed and extrapolated OS estimates from multiple data cuts using standard parametric and spline-based approaches

Data

- This analysis compares observed and extrapolated OS estimates from 3 data cuts from Part A of the JAVELIN Merkel 200 trial
- Each data cut constitutes a different period of minimum follow-up (MFU) for all patients:
 - Data cut Sep-2016
 - 12 months' MFU
 - Data cut Mar-2017
 - 18 months' MFU
 - Data cut lock Sep-2017
 - 24 months' MFU

Kaplan-Meier plot of OS for each data cut



	Sep '16	Mar '17	Sep '17
Sep '16	88	59	41
Mar '17	88	59	42
Sep '17	88	60	42

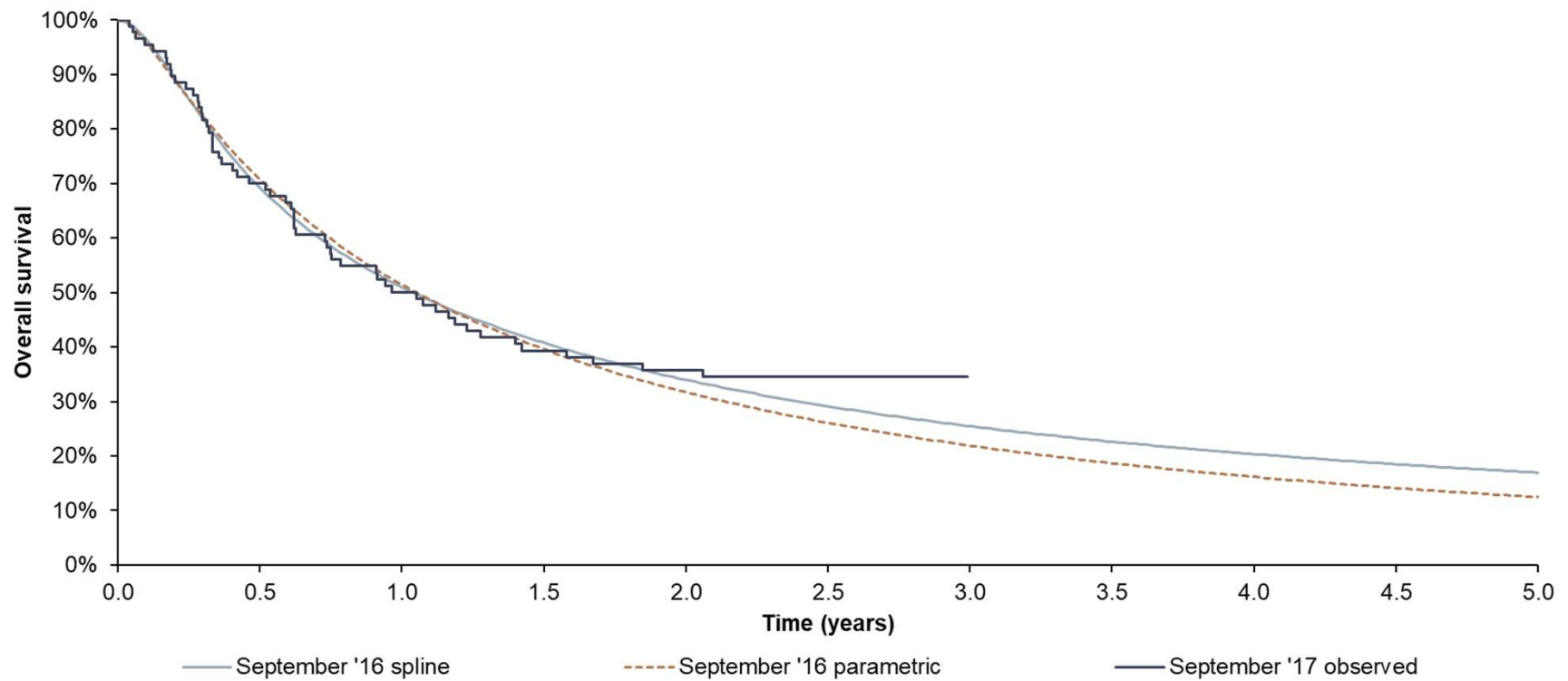
12 months' MFU: Kaufman *et al.*, (2018) <https://www.ncbi.nlm.nih.gov/pubmed/29347993>
 18 months' MFU: D'Angelo *et al.*, (2018) http://ascopubs.org/doi/abs/10.1200/JCO.2018.36.5_suppl.192
 24 months' MFU: Nghiem *et al.*, (2018) <https://meetinglibrary.asco.org/record/161628/abstract>

Methods

- Standard parametric and spline-based models were fitted to OS data from Part A of JAVELIN Merkel 200 for each data cut
- All survival models were fitted in the statistical software R using the flexsurv package
 - The standard parametric survival models considered were the exponential, generalised gamma, Gompertz, log-logistic, log-normal, and Weibull (routinely considered to inform the estimation of OS in health technology assessment)
 - The spline-based models considered were natural restricted cubic spline models. The spline-based models were fitted with 1-3 internal knots using each of the 3 functional forms permitted by flexsurv. Knot locations were selected according to the percentiles of the log-uncensored survival times
- The selection of the best-fitting parametric survival model was determined through a combination of visual fit to the observed OS data, statistical goodness-of-fit (measured by Akaike's information criterion [AIC]), and the plausibility of long-term extrapolation (based on clinical expert input)

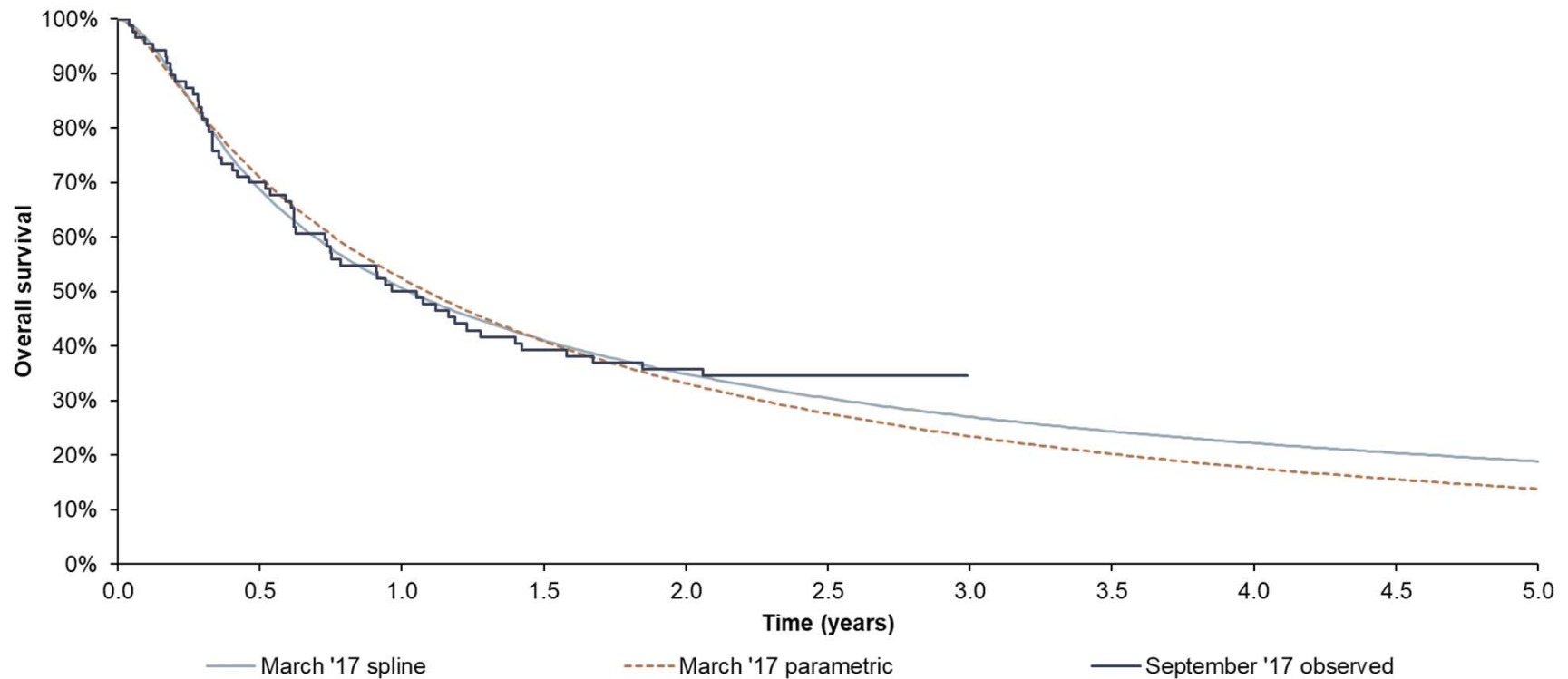
Results

Extrapolation from September '16 (12 months' MFU) data cut



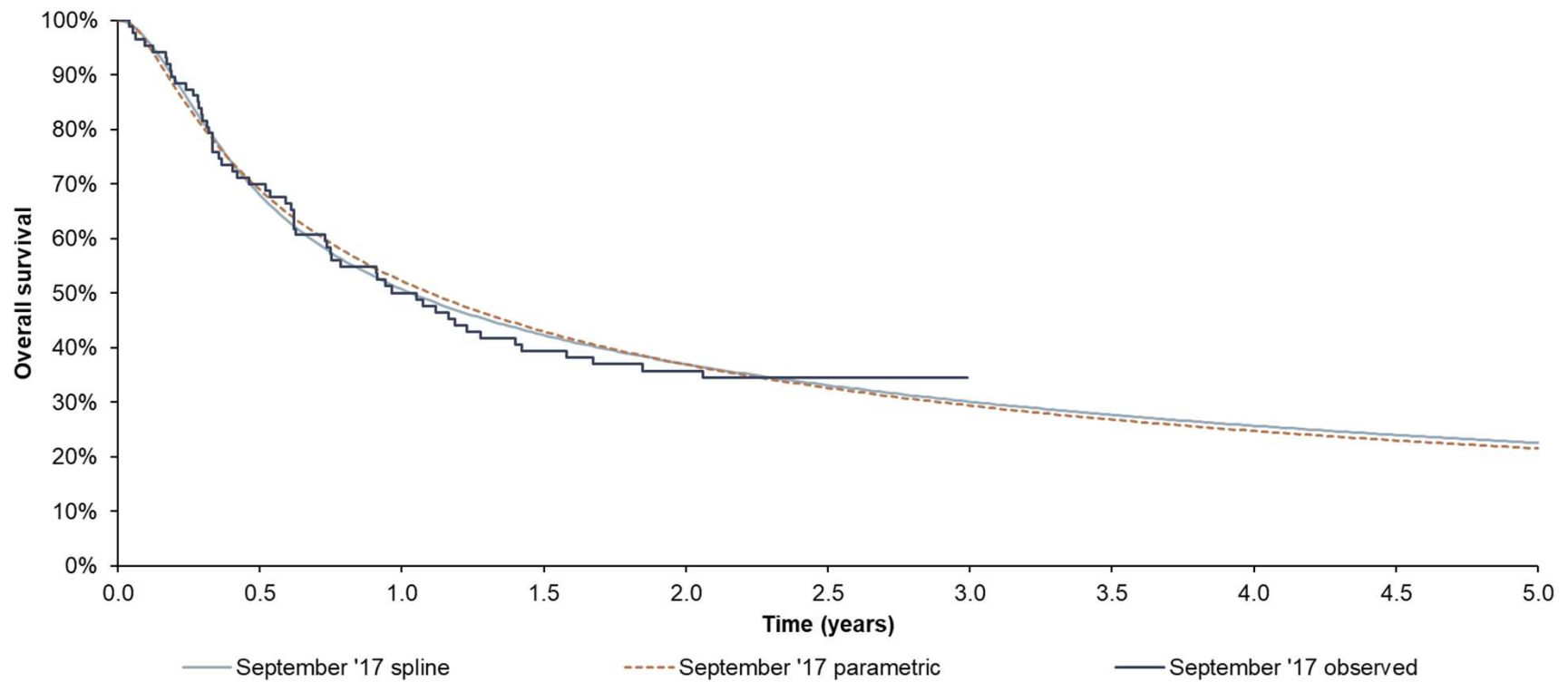
Results

Extrapolation from March '17 (18 months' MFU) data cut



Results

Extrapolation from September '17 (24 months' MFU) data cut



Results Summary

Assessment time			12 months	18 months	24 months	
Observed survival						
September 2016 (12 months' MFU) data cut			51.8%	38.3%	38.3%	
March 2017 (18 months' MFU) data cut			50.8%	39.9%	34.3%	
September 2017 (24 months' MFU) data cut			50.1%	39.3%	35.8%	
Assessment time	Model	AIC	12 months	18 months	24 months	60 months
Standard parametric survival						
September 2016 (12 months' MFU) data cut	Log-normal	377.70	51.5%	39.3%	31.8%	12.4%
March 2017 (18 months' MFU) data cut	Log-normal	431.54	52.5%	40.7%	33.3%	13.8%
September 2017 (24 months' MFU) data cut	Log-normal	455.31	54.1%	42.9%	35.8%	16.2%
	Generalised gamma	454.68	52.3%	42.8%	37.0%	21.5%
Spline-based survival						
September 2016 (12 months' MFU) data cut	1-knot odds	379.26	51.5%	40.5%	34.1%	16.9%
March 2017 (18 months' MFU) data cut	1-knot odds	432.29	50.7%	40.9%	35.0%	18.9%
September 2017 (24 months' MFU) data cut	1-knot odds	453.81	50.8%	42.1%	37.1%	22.5%

Conclusions

- Spline-based models provided a more accurate estimation of the observed 24-month OS based on extrapolation from earlier data than standard parametric approaches
- Longer-term survival estimates from the spline-based models were more aligned with clinical expectations of immunotherapy, ie, an emergent plateau in OS associated with the immune-response effect of treatment
- Limitations and further research:
 - Landmark or cure-based models may also reflect the expected immune-response effect in OS but require explicit assumptions about the estimation of long-term OS (such as the OS for cured patients, the prognostic importance of response, and the difference in the hazard of death by response)
 - Longer-term data are required to validate OS extrapolations

Thank you

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