



**Food and Drug Administration Advisory Committee Member
Acknowledgment of Financial Interests**

Name of Advisory Committee Member: **Colin Weekes, MD, PhD**

Committee: **Oncologic Drugs Advisory Committee (ODAC)**

Meeting Date: **April 29, 2021**

I acknowledge that contingent upon public disclosure of the following financial interests related to the agenda items described below, I may be considered for participation in the advisory committee meeting.

On April 29th, the committee will hear updates on (1) BLA 125514/S-024, trade name Keytruda (pembrolizumab), submitted by Merck Sharp & Dohme Corp. and indicated for the treatment of patients with recurrent locally advanced or metastatic gastric or gastroesophageal junction adenocarcinoma whose tumors express PD-L1 [Combined Positive Score (CPS) ≥ 1] as determined by an FDA-approved test, with disease progression on or after two or more prior lines of therapy including fluoropyrimidine and platinum-containing chemotherapy and if appropriate, HER2/neu-targeted therapy; (2) BLA 125514/S-042, trade name Keytruda (pembrolizumab), submitted by Merck Sharp & Dohme Corp. and indicated for the treatment of patients with hepatocellular carcinoma (HCC) who have been previously treated with sorafenib; and (3) BLA 125554/S-041, trade name Opdivo (nivolumab), submitted by Bristol Myers Squibb Company and indicated as a single agent for the treatment of patients with hepatocellular carcinoma (HCC) who have been previously treated with sorafenib.

These applications were approved under 21 CFR 601.40 (subpart E, accelerated approval regulations) with confirmatory trial(s) that have not verified clinical benefit. These updates will provide information on: (1) the status and results of confirmatory clinical studies for a given indication; and (2) any ongoing and planned trials. Confirmatory studies are post-marketing studies to verify and describe the clinical benefit of a drug after it receives accelerated approval. Based on the updates provided, the committee will have a general discussion focused on next steps for each product including whether the indications should remain on the market while additional trial(s) are conducted.

<u>Type of Interest</u>	<u>Nature</u>	<u>Magnitude</u>
I. Personal/Immediate Family		
None		
II. Other Imputed Interests		
Contracts/grants	Employer's research funded by Bristol Myers Squibb, sponsor of Opdivo (nivolumab)	\$50,000 – \$100,000, per year

Contracts/grants	Employer's anticipated research funded by (b) (4)	(b) (4), per year anticipated from (b) (4) \$ (b) (4) per year, in salary support
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I hereby request that FDA make this information publicly available on my behalf if the agency grants a waiver allowing me to participate in the meeting described above. I understand that without public disclosure of these interests, I will not participate in the advisory committee meeting described above.

_____/S/_____
Signature

_____/4/1/2021_____
Date