Exposure–Response Analysis of Overall Survival for Tremelimumab in Unresectable Malignant Mesothelioma: The Confounding Effect of Disease Status

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Tremelimumab, an anti-cytotoxic T-lymphocyte antigen-4 monoclonal antibody that enhances T-cell activation, was evaluated in a randomized, double-blind, placebo-controlled, phase IIb study (NCT01843374) in patients with unresectable malignant mesothelioma. The study demonstrated no clinically meaningful differences in overall survival (OS). The objective of this analysis was to evaluate the relationship of exposure with OS. A population pharmacokinetic (PK) model adequately described the PK data. Three factors (sex, C-reactive protein, and baseline tumor size) were identified as statistically significant PK predictors (P < 0.05 on clearance). A positive association between exposure and OS was observed. However, an association between key baseline factors with OS (regardless of treatment) and imbalances in prognostic factors favoring patients with higher exposure (upper vs. lower PK quartile) was seen. Taken together, these results suggest that the exposure OS relationship observed for tremelimumab in mesothelioma is likely spurious rather than a true association of exposure with efficacy.

Tremelimumab is a human monoclonal antibody that binds to CTLA-4 and blocks its interaction with its ligands. Tremelimumab single-agent was investigated in melanoma and in unresectable pleural and peritoneal malignant mesothelioma but did not provide sufficient efficacy to warrant approval. However, tremelimumab is currently investigated in a number of malignancies in combination with another checkpoint monoclonal antibody inhibitor, durvalumab, which targets programmed cell death ligand 1.

Study Highlights

WHAT IS THE CURRENT KNOWLEDGE ON THE TOPIC? ▶ Tremelimumab is a human monoclonal antibody that binds to CTLA-4 and blocks its interaction with its ligands. Tremelimumab single-agent was investigated in melanoma and in unresectable pleural and peritoneal malignant mesothelioma but did not provide sufficient efficacy to warrant approval. However, tremelimumab is currently investigated in a number of malignancies in combination with another checkpoint monoclonal antibody inhibitor, durvalumab, which targets programmed cell death ligand 1.

WHAT QUESTION DID THIS STUDY ADDRESS? ▶ This work presents a post hoc analysis evaluating the relationship of tremelimumab exposure with overall survival (OS). In addition, an analysis to further evaluate the potential association among disease factors, exposure, and OS was performed.

WHAT DOES THIS STUDY ADD TO OUR KNOWLEDGE? ▶ The OS exposure–response relationship for tremelimumab has not yet been described in malignant mesothelioma. This study proposes a pragmatic but systematic approach to data analysis, coupled with deductive reasoning, to decipher the OS exposure response through integrated analysis of trial data, including risk factors.

HOW MIGHT THIS CHANGE CLINICAL PHARMACOLOGY OR TRANSLATIONAL SCIENCES? ▶ A simplistic empirical approach to exposure–response analysis, when complemented by a review of baseline factors associated with OS, population pharmacokinetic analysis and review of baseline factors across exposure groups, can delineate the multivariate factors underlying an apparent trend in OS exposure response. Our pragmatic analysis provides an alternative approach to case control, especially in oncology, where generally no adequately powered dose-response-finding trials with comparator arm can be conducted.
to CTLA-4 by anti-CTLA-4 antibodies results in markedly enhanced T-cell activation and antitumor activity in animal models, including killing of established murine solid tumors and induction of protective antitumor immunity. Therefore, it is expected that treatment with tremelimumab will lead to activation of the human immune system, increasing antitumor immunity in patients with solid tumors.

Although phase II and phase III studies of tremelimumab in metastatic melanoma did not meet the primary end points of response rate and overall survival (OS), respectively, the data suggested clinical activity. In a large phase III randomized study (NCT00257205), tremelimumab 15 mg/kg administered by i.v. infusion every 12 weeks (q12w) failed to demonstrate survival benefit compared with first-line standard of care treatment with a reported median OS of 12.6 months for tremelimumab vs. 10.7 months for chemotherapy (hazard ratio = 0.88, P = 0.127). Opportunistically, mesothelioma was considered to have an unmet medical need that could potentially be addressed by anti-CTLA-4 therapy. This was motivated by a promising sign of efficacy from two phase II investigator-sponsored studies evaluating tremelimumab in second-line unresectable metastatic malignant mesothelioma.9,10 Mesothelioma is a lethal disease that has one of the worse prognoses among solid tumors, with < 5% of patients surviving 5 years. The estimated median survival time in untreated cases ranged from 6–9 months from the date of diagnosis.11 For advanced disease, cisplatin and pemetrexed combination therapy is standard of care for first-line treatment for pleural mesothelioma, with an approximate 3-month increase in median OS in patients treated with a platinum-based regimen. Patients were randomized (2:1) to receive either tremelimumab (n = 382) at 10 mg/kg q4w (seven doses) followed by q12w i.v. or placebo (n = 189). However, the study demonstrated no clinically meaningful differences in OS12,13 and motivated an in-depth analysis, including tremelimumab pharmacokinetics (PK) and its relationship with efficacy. Currently, tremelimumab is investigated in various indications in combination with another checkpoint monoclonal antibody inhibitor, durvalumab, that targets programmed cell death ligand 1.

Tremelimumab PK properties were previously reported based on a population PK modeling approach combining data from phase I, II, and III studies (n = 654) in subjects with metastatic melanoma using nonlinear mixed-effects modeling.14 A two-compartment linear PK model, consistent with a natural immunoglobulin G2 molecule, adequately described the plasma concentrations of tremelimumab following various dosing regimens. The population estimates for clearance (CL) and central volume of distribution (V1) were 0.26 L/day and 3.97 L, respectively, with modest interindividual variability (31.8% and 20.4%, respectively). CL was higher in men, subjects with higher values of creatinine clearance and endogenous immunoglobulin G, and subjects with relatively poor baseline prognostic factors: Eastern Cooperative Oncology Group (ECOG) status (higher CL for ECOG > 0), lactate dehydrogenase (LDH) levels (higher LDH resulted in higher CL), and C-reactive protein (CRP) levels (higher CRP resulted in higher CL). Central volume of distribution was higher in men and subjects with higher body weight. No dose adjustment was needed based on the magnitude of the change in PK. Similar to other monoclonal antibodies without target-mediated drug disposition, tremelimumab is likely to be cleared from circulation by endothelial cell uptake and proteolysis.15 Hence, no impact of renal/hepatic functions is expected on tremelimumab elimination.

The same model structure fitted equally well the PK data from the two investigator-initiated studies (NCT01649024/NCT01655888) in 40 patients with malignant mesothelioma (18 patients of the pooled analysis did not have PK data), confirming similar PK properties of tremelimumab across the two indications (CL and V1 were 0.2 L/day and 3.5 L, respectively, with body weight and ECOG performance status impacting exposure levels).

The primary objective of this analysis was to provide a systematic evaluation of the relationship of PK exposure...
with OS of the DETERMINE trial to demonstrate that tremelimumab dosing regimen fully tested the mechanism of action in patients with mesothelioma. Specifically, we aimed to derive individual PK metrics by PK modeling to evaluate potential relationship between exposure and OS. Subsequently, to further test any observed E-R relationship, we assessed any potential factors other than exposure on OS and investigated any imbalance of disease status in each exposure subset. The statistical significance of potential confounders found in E-R investigation was then tested by population PK covariate analysis.

METHODS

The methodology applied in this work was performed in a three-step process that involved empirical evaluation through graphical analysis of OS data and the use of population PK analysis to confirm determinants of tremelimumab PK. The first step consisted of obtaining a reliable representation of steady-state exposure levels for each patient who received at least one dose of tremelimumab in the DETERMINE trial. The intent was to obtain a standardized PK metric to evaluate the degree of association of PK with OS data from this trial. In order to circumvent the limitations of observed concentration data (data handling issues associated with missing data, PK data obtained prior to achieving steady state for some patients, measurement errors, and other sources of residual variability in exposure measurements), use of model-derived PK exposure metrics, such as the area under the exposure time-course curve at steady-state \( \text{AUC}_{ss} \) was considered as a robust and unbiased approach. To this aim, a population PK model of tremelimumab was developed based on the DETERMINE trial observed PK data and was used to predict steady-state exposure metrics for each patient.

The second step of this analysis consisted of a graphical evaluation using a Kaplan–Meier (KM) plot of OS split by tremelimumab exposure levels. \( \text{AUC}_{ss} \) were ranked into quartiles and used to identify any potential underlying relationship between steady-state PK exposure and OS data. An analysis was then performed using KM plots of OS split by baseline disease status in both the treatment group and the control group to evaluate any trend on the OS profile suggestive of potential prognostic or predictive factors. Given the presence of time-dependent predictive factors (i.e., exposure) and the degree of collinearity between predictors, which are likely to confound interpretation, a multivariate Cox regression model has not been conducted.

The last step of the analysis consisted of evaluating a potential correlation between disease status and PK exposure in an attempt to identify any important factors that may explain the relationship found in step 2 of this analysis. This was performed by means of covariate analysis of the population PK model.

**Step 1: Base population PK model development**

Population PK of tremelimumab was based on nonlinear mixed-effects modeling methodology, implemented in the computer program NONMEM, version 7.3.\(^{16}\) Model development started with the base model (i.e., the best description of the data without considering the effect of covariates). The modeling details are provided in **Supplementary Materials S1**.

**Step 2: Exposure–efficacy (OS) analysis and confounding analysis**

Once a base model had been established, individualized PK exposure metrics were estimated (specifically, \( \text{AUC}_{ss} \) and CL) for E-R analysis of OS data. An exposure–OS relationship was evaluated by a KM plot of OS by quartiles of the \( \text{AUC}_{ss} \) distribution performed in R (version 3.3.1 or higher\(^{17}\)).

A similar exploratory evaluation on OS data was done based on several risk factors measured at baseline. Instead of using exposure as a potential predictor of OS, as is typically done for E-R analysis, the KM plot of OS was presented based on patients’ characteristics at baseline. A pool of 13 potential baseline confounders was defined a priori based on mechanistic plausibility, scientific interest, and prior knowledge. It consisted of the three stratification factors of the trial that are considered prognostic in mesothelioma (European Organisation for Research and Treatment of Cancer (EORTC) status, anatomic site, and line of therapy), as well as the additional factors of histology, ECOG performance status, race, sex, age, body weight, inflammatory status as measured by CRP and serum albumin, LDH levels, and baseline tumor burden. Categorical covariates were split based on subgroups defined in Maio et al.\(^{13}\) For continuous covariates, the median was used as a cutoff to dichotomize patients into high or low levels. Because the DETERMINE trial included a placebo comparator arm, the KM plot of OS was split by treatment arm (tremelimumab and placebo) for each of the 13 risk factors to evaluate their prognostic/predictive value. Risk factors for which a trend was visible on visual exploration of survival were then evaluated for potential imbalance in each \( \text{AUC}_{ss} \) exposure quartile.

**Step 3: Covariate analysis to confirm potential PK predictors**

After completion of base population PK model development, each of the 13 covariates (evaluated in step 2 as potential risk factors on OS) was tested for their explanatory power on PK model parameters by means of likelihood ratio statistical testing with a type-I error of 5%. The effect of antidrug antibodies (ADAs) on tremelimumab exposure was also examined. The clinical or physiological relevance for each covariate effect was ultimately evaluated for its significance to establish the final population PK model. More details are provided in **Supplementary Materials S1**.

Finally, a presentation of the multidimensional relationship among PK, disease factors, and OS was attempted using the iGraph package version 1.1.2 in R.\(^{17}\)

**RESULTS**

**PK modeling**

The PK analysis data set from the DETERMINE study included 376 patients with mesothelioma and 1,328 evaluable PK concentrations (six patients had no PK evaluable data). Tremelimumab PK (Table 1) was consistent with previous reports. A two-compartment linear model adequately
Given that the population model adequately described the PK of tremelimumab and that no covariate was found to be clinically relevant, the final PK model was equivalent to the base model (with no covariate) used to derive individual PK metrics (AUC<sub>ss</sub>). The distribution of predicted AUC<sub>ss</sub> values representative of individuals’ exposure level used for investigation of E-R based on observed OS is displayed in Figure S2.

### Exposure–OS analysis

Similar OS data were observed for tremelimumab (median OS time = 7.72 months) compared with placebo (7.29 months) based on the intention-to-treat population (Figure 1 left panel). The hazard ratio from stratified Cox regression model (including the two stratification factors EORTC status and line of therapy) was 0.92 (95% CI: 0.76–1.12, P = 0.408).<sup>12,13</sup>

When assessed, based on tremelimumab PK exposure (AUC<sub>ss</sub>), a monotonic relationship with OS was observed, with the all-comers curves centered between the interquartile range (Q2 and Q3; Figure 1 right panel and Table 3). The median survival in the tremelimumab group for patients with the highest quartile of exposure was 14.9 months (95% CI: 12.5–18.6). A twofold increase in median AUC<sub>ss </sub>exposure level between the most extreme exposure quartiles (Q1 and Q4) resulted in a threefold improvement in median OS time (Table 3). However, although increasing exposure seemed to yield a better OS profile, the lowest AUC<sub>ss </sub>quartile (Q1) median survival was worse than placebo (4.93 vs. 7.29 months), with its 95% CI (3.86–6.68) not including the placebo point estimate; this suggests that some confounders are likely affecting the E-R relationship. A sensitivity analysis based on observed peak plasma concentration (C<sub>max</sub>) after the first cycle of treatment led to a similar trend (Figure S3).

### Table 1 Final population PK model parameter estimates of tremelimumab from the DETERMINE trial

| Model parameter | θ (Median) | RSE (θ) (%) | BSV (Ω) (%) | RSE (Ω) (%) |
|-----------------|------------|-------------|-------------|-------------|
| CL (L/day)      | 0.310      | 4.62        | 38.0        | 8.54        |
| V1 (L)          | 3.85       | 1.91        | 32.5        | 16.5        |
| V2 (L)          | 1.72       | 19.4        | 25.2        | 84.6        |
| Q (L/day)       | 0.273      | 42.8        |             |             |
| CV%             | 37.7%      | 4.92        |             |             |

BSV, between-subject variability of parameter with random effect assumed normally-distributed with mean θ and variance Ω<sup>2</sup>; CL, clearance; CV%, coefficient of variation percentage; PK, pharmacokinetic; RSE, relative standard error obtained from $COVARIANCE$ step in NONMEM; V, volume of distribution.

described the data. CL and V1 estimates from the final PK model were 310 mL/day and 3.85 L, with moderate variability of 38.0% and 32.5%, respectively, vs. 260 mL/day (31.8% between-subject variability) and 3.97 L (20.4%) in melanoma.<sup>10</sup> Fifteen (4%) of 377 patients with ADA evaluable data were ADA-positive postbaseline in the treatment arm, which was comparable to the placebo arm (3.2%, 6% associated with significantly lower tremelimumab exposure levels (< 6%). ADA had no apparent effect on PK.

Higher baseline tumor size, higher CRP levels, and men were associated with significantly lower tremelimumab exposure levels (P < 0.05; Table 2). These results were consistent with previous findings<sup>10</sup> but were not found clinically relevant with respect to differences in exposure (< 30% effect on AUC<sub>ss</sub>; Figure S1).

### Table 2 Population PK covariate analysis of tremelimumab from the DETERMINE trial

| Hierarchical model | Covariate-PK relationship | Covariate-PK relationship strength | OFV likelihood | Reference OFV | P value |
|--------------------|---------------------------|-----------------------------------|----------------|--------------|---------|
| 1. Base model      | None                      | –                                 | 10,428         | –            | –       |
| 2. Intermediate model 1 | CL % reduction for female patients vs. male | –0.19 | 10,414 (Δ = −14) | 1 | 0.00018 |
| 3. Intermediate model 2 | Serum albumin on CL | 0.00094 | 10,413 (Δ = −1) | 2 | 0.32 |
| 4. Intermediate model 3 | CRP effect on CL | 0.0034 | 10,370 (Δ = −44) | 2 | <10<sup>−10</sup> |
| 5. Intermediate model 4 | ECOG effect on CL | 0.019 | 10,368 (Δ = −1.4) | 4 | 0.24 |
| 6. Intermediate model 5 | EORTC effect on CL | 0.068 | 10,368 (Δ = −2.0) | 4 | 0.16 |
| 8. Intermediate model 6 | LDH effect on CL | −2.1 × 10<sup>−04</sup> | 10,364 (Δ = −1.3) | 7 | 0.26 |
| 5. Intermediate model 7 | Histology (epithelioid) % increase on CL | 0.084 | 10,363 (Δ = −1.7) | 7 | 0.19 |
| 7. Final model | Baseline tumor size effect on CL | 0.00058 | 10,365 (Δ = −4.3) | 4 | 0.039 |

Continuous covariates were entered in the model assuming proportional linear effect of the covariate on PK from median cutoff (median is 31 g/L for albumin, 33 mg/L for CRP, and 97 mm for baseline tumor size). Sex, CRP, and baseline tumor size were significant PK predictors and explained 20% of the individual variability on CL, reducing coefficient of variation from 42% to 38%. ΔOFV was computed for nested models according to the order displayed in the “Reference OFV” column, where the reference OFV is taken as the previous model with the statistically significant covariate-PK relationship included. CL, clearance; CRP, C-reactive protein; ECOG, Eastern Cooperative Oncology Group; EORTC, European Organisation for Research and Treatment of Cancer; LDH, lactate dehydrogenase; OFV, objective function value; PK, pharmacokinetic.
Potential confounders of the tremelimumab E-R in OS of patients with mesothelioma were assessed by evaluating imbalance of patients' characteristics in the extreme quartiles of exposure. Analysis of OS data based on baseline patient characteristics other than exposure revealed a number of factors as important, regardless of treatment. Higher OS was observed in patients with lower CRP levels, in patients with ECOG performance status = 0 compared with ECOG = 1, and in patients with EORTC low risk compared with EORTC high risk (Figure 2). Specifically, when assessing by performance status across treatment arms, OS data show ECOG = 0 patients survive longer than ECOG = 1 patients (median OS = 11–13 vs. 5–7 months) irrespective of treatment received. Review of baseline characteristics across AUC groups indicated that there were more ECOG = 0 patients in Q4 of AUCss (highest tremelimumab exposure) than Q1 (lowest exposure; Figure 3). Longer OS was also observed in patients with low-risk (ECOG = 1) compared with high-risk status (EORTC = 2), with a median OS by EORTC status across treatment groups of 10 vs. 6 months, respectively (Figure 2). Conversely, an imbalanced distribution of low-risk/high-risk EORTC patients was visible across the exposure quartiles, with more low risk (as assessed by EORTC) patients in Q4 (highest exposure) than Q1 (lowest exposure) (Figure 3). Similarly, data suggested that high tumor burden (tumor size at baseline ≥ median = 97 mm) was associated with shorter survival (median OS = 11–9 vs. 7 months), although this is more visible in the tremelimumab treatment group (Figure 2).

Patients with high tumor burden at baseline demonstrated lower PK exposure (Q1 of AUCss; Figure 3). Longer OS was observed in patients with lower inflammatory biomarker levels (CRP < median = 33 mg/L; i.e., ~11-fold the upper limit of normal) compared with patients with higher inflammatory biomarker levels (Figure 2). Patients with high CRP were more common in the lower PK exposure group (Q1 of AUCss; Figure 3). Last, longer OS was observed in women than in men, albeit the tails of the OS KM plot converge at the later timepoints (Figure 2). This finding is aligned with a recent publication that found women had a 28% lower mortality rate than men in pleural mesothelioma.18 Men had higher CL than women (Q4 of CL) and, thus, had lower PK exposure (Q1 of AUCss; Figure 3).

Based on these findings, a map of risk factors identified for OS and of the covariates impacting PK was built (Figure 4). This map was adapted based on the original mapping of confounding factors by Skelly et al.19

**DISCUSSION**

The understanding of confounding factors in E-R analysis is evolving in oncology.20,21 This is primarily motivated by two factors: because of trial design issues specific to oncology, PK information is usually limited and encompassing a narrow dose–response range; and, especially for biotherapeutics, cancer progression and patient health status may affect PK.22,23 This exacerbates the issue of having to decide whether observed differences in OS...
between standard of care and experimental treatment are due to imbalances in confounding risk factors at baseline or differences in drug exposure. When a single-dose experimental arm fails to demonstrate superiority over standard of care (as in the tremelimumab confirmatory trial in melanoma), post hoc analysis of E-R is generally conducted to delineate whether a higher dose could have resulted in survival benefit, provided that the maximum tolerated dose was not yet tested. When the single-dose experimental arm shows a clinically relevant benefit over standard of care, as in the case of trastuzumab in gastric cancer or trastuzumab-DM1 in breast cancer, a post hoc analysis of E-R was also conducted to decide whether nonresponders could have benefited from a higher dose. In both cases, this can result in further confirmatory trials with a higher dose (as in tremelimumab in mesothelioma) or in postapproval commitment trials, as exemplified by the HELOISE trial evaluating higher dose of trastuzumab and by the TH3RESA trial for trastuzumab-DM1. For both tremelimumab and trastuzumab, results demonstrated no additional benefit of higher doses and, thus, put into question the conclusions drawn from such E-R post hoc analyses.

Case-control E-R analysis has been suggested to ascertain whether baseline risk factors known to affect OS and associated with drug exposure have a role in changes in OS. Case-matched control comparison allows for the definition of matched subgroups that minimize differences in patient characteristics, thus resulting in balanced distributions of measured risk factors that allow the role of differences in exposure to be ascertained. However, this methodology may lack robustness, as demonstrated by the negative outcome of the HELOISE trial that was predicated from such an analysis. One possible reason is that it does not account for the dynamic nature of the intricate interactions among risk factors, survival, and exposure, which can change relatively rapidly with time, especially when PK is reaching steady state (the most relevant exposure metric when conducting E-R). As a workaround, Wang et al. suggests to use an early metric of exposure (e.g., AUC at the end of the first cycle). The reasoning behind this is that early assessment of PK metrics may be seen as less affected by longitudinal changes in therapeutic antibody clearance over time driven by disease status. However, the effect of disease status on PK at baseline is not considered, nor the change of exposure within individuals over time; therefore, more sophisticated approaches that account for longitudinal changes in both disease markers and PK are preferable.

The E-R post hoc analysis of OS data from the DETERMINE study suggested a trend in patients exposed to tremelimumab, with higher exposure levels resulting in longer survival. Specifically, this translated into the highest quartile of the AUC group with a reasonable sample size (n = 94) showing a large difference of ~8 months in median OS over the all-comers control arm (n = 189). Such effect size could be considered a breakthrough therapy.
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for unresectable malignant mesothelioma, a disease with a dismal outcome and no approved treatment option beyond first-line therapy. Conversely, the lowest quartile of the AUC\textsubscript{ss} subgroup did worse than the all-comers control arm (4.9 vs. 7.3 months median survival). This raised suspicion about a confounding bias induced by the E-R categorization because no biologically plausible hypothesis could be proposed for why low tremelimumab exposure would cause a worse survival outcome than placebo. A sensitivity analysis using an early metric of exposure found similar E-R as observed when using steady-state PK. Instead of conducting a case-control E-R analysis that would have further reduced the sample size, potential confounding factors were initially explored to assess their impact. Through graphical evaluation of OS using KM plots, we identified five potential risk factors impacting OS: sex, ECOG performance status, EORTC prognostic status, inflammatory status (CRP), and tumor burden at baseline. Based on the biology of malignant tumors, these factors are expected to relate to OS regardless of tremelimumab exposure, and any imbalance in these baseline factors when grouping by exposure status may result in a confounding effect, making the E-R only “apparent.” Interestingly, in the DETERMINE trial, patients in the highest exposure quartile (Q4 AUC\textsubscript{ss} or Q1 CL) exhibited more favorable prognostic factors (low CRP, female, low tumor burden, ECOG = 0, and low-risk EORTC) compared with patients in the lowest exposure quartile (Q1 AUC\textsubscript{ss} or Q4 CL). Likewise, more patients with poor prognostic factors (high CRP, male, high tumor burden, ECOG = 1, and high-risk EORTC) were belonging to the lowest exposure quartile than to the highest exposure quartile.

Subsequently, we used nonlinear mixed-effects modeling of the DETERMINE trial of tremelimumab longitudinal PK data to decipher whether any of these five baseline disease factors could also affect tremelimumab exposure. Conceptually, if an interaction between PK and a known risk factor of OS was identified, this could translate into an indirect relationship between PK and OS, the resultant being an apparent trend of E-R when this is explored in a simplified framework, such as KM plots. Our analysis found that at least three baseline factors (sex, CRP, and baseline tumor size) were statistically significant predictors of tremelimumab PK (P < 0.05 for CL), indicating a multidimensional confounding effect. Thus, the observed apparent exposure–OS relationship of tremelimumab is likely due to the imbalance in key baseline factors across the tremelimumab PK groups and the association of baseline prognostic factors for OS with PK (CL), rather than a true association of exposure with efficacy.
In conclusion, this analysis supports that patients with higher tremelimumab exposure did not derive additional benefit from this treatment after chemotherapy, despite signs of biological activity of anti-CTLA-4 therapy as judged by the higher incidence of immune-mediated adverse events (e.g., colitis) observed in tremelimumab-treated patients. In light of the confounding risk factors affecting OS and the correlation of some risk factors with PK, we conclude that the apparent E-R relationship observed for tremelimumab in mesothelioma is spurious. In contrast, the standard assumption used in E-R analysis (i.e., considering PK as an independent predictor variable of the dependent outcome variable OS) is invalidated by the imbalance of risk factors across exposure groups and relationship of risk factors with both PK and OS. The deductive map shown in Figure 4 provides a simplistic view to cogently distinguish apparent proximate cause (PK) from root causes (disease risk factors) that govern mortality in mesothelioma. This implies that careful consideration should be given not to over-interpret empirical E-R results, and complementary analyses, such as dose–response and mixed-effects modeling, should be relied upon to prevent misleading conclusions.

Supporting Information. Supplementary information accompanies this paper on the Clinical and Translational Science website (www.cts-journal.com).

Figure S1. Effect of baseline covariates on exposure parameter $\text{AUC}_{\text{ss}}$. Figure S2. Distribution of tremelimumab predicted exposure at steady-state ($\text{AUC}_{\text{ss}}$) in all-comers and across each quartile ($Q_1 =$ lowest exposure, $Q_3 - Q_2 =$ interquartile range, and $Q_4 =$ highest exposure).

Figure S3. Kaplan–Meier sensitivity analysis of OS by quartiles of observed exposure levels after the end of infusion of the first dose of tremelimumab ($C_{\text{max,1}}$) for the tremelimumab-treated group.

Supplementary Material S1. Methods.

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