# A Description of the IVI-NSCLC Model v1.0 $\,$

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# 1 Open Source Value Project

The continuing increase in U.S. healthcare costs has stimulated the introduction of initiatives to promote the use of high-value care. Cost-effectiveness analysis can inform efficient use of healthcare resources by formally computing costs and benefits to identify the most valuable treatment options for a given disease. In many countries, a single health technology assessment agency assesses the value of healthcare technology by means of cost-effectiveness analysis and recommends a utilization strategy. In the US, however, utilization decisions are decentralized and made by a variety of payers and provider organizations. Value frameworks are gaining prominence to guide utilization of therapies, but vary in perspective, the evidence considered, and approaches, thereby resulting in confusion and debate among stakeholders.

A thorough evidence-based analysis of the value of medical technology is resource intensive and complex. Typically, there is no empirical study with sufficient long-term follow-up that compares all treatments for a particular disease regarding relevant clinical outcomes and costs. Thus, costeffectiveness analyses generally rely on mathematical models that integrate evidence on the course of disease, treatment effects, and the relationship between clinical outcomes and costs, from a variety of studies. The nature of these evaluations can lead to disputes in the scientific literature and community. Models are typically difficult to understand. Even modeling experts may not be able to fully understand a model-based cost-effectiveness analysis without public source code and detailed model documentation. This lack of transparency also poses problems for users whose perspective, local context, or patient population varies from that of the reported analysis. In the absence of public access to the actual model, updating the evaluation is cumbersome, if not impossible, for someone other than the original model developer. As a result, published cost-effectiveness findings risk immediate irrelevance to some stakeholders and growing irrelevance to all stakeholders as new clinical evidence emerges. Value assessment only has relevance for decision-making when it reflects the totality of the latest evidence, is transparent, deemed credible by different stakeholders, representative of the local context and patient population, and can be easily updated without duplication of effort.

With the Open-Source Value Project (OSVP), the Innovation and Value Initiative (IVI) aims to maximize both the relevance and credibility of value assessment in the context of the United States' decentralized decision-making environment by developing and providing access to flexible open-source decision models for value assessment. These interactive models have two primary objectives: (i) to enable a more constructive dialogue regarding value assessment between stakeholders (e.g. patients, payers, providers, and manufacturers) with different beliefs about relevant clinical data, modeling approaches, and value perspectives; and (ii) to provide local decision-makers with means to credible value assessment that reflects the local setting and is based on the latest evidence while

accounting for scientific uncertainty (due to patient heterogeneity, gaps in evidence, and different modeling beliefs).

In order for a decision model to remain relevant over time it needs to evolve along with the clinical evidence and scientific insights. OSVP facilitates iterative development and collaboration between multiple clinical and methodological experts. Refer to the IVI website for additional information about the OSVP.

# 2 Topic definition

IVI was interested in developing its most recent OSVP within oncology. When selecting the tumor type of interest, IVI considered criteria such as burden of disease, development of innovative treatments, alternative treatment strategies available, availability of clinical evidence, and engagement of patient organization(s) to actively contribute to the project.

Lung cancer is the leading cause of cancer related death worldwide (Jemal et al. 2011). Non-small lung cancer (NSCLC) accounts for an estimated 85% of lung cancer cases and comprises adenocarcinoma, squamous cell carcinoma, and large cell carcinoma (D'addario et al. 2010). The five-year survival of stage IV NSCLC is less than 2% (Cetin et al. 2011). Given the rapid pace of development of new therapies in NSCLC, the scope of the OSVP needed to be limited to a specific sub-population in order to ensure a model could be developed in a reasonable time frame in a manner consistent with the evidence base. The selected target population of interest for the most recent OSVP is metastatic epidermal growth factor receptor positive (EGFR+) NSCLC. EGFR mutations are more commonly observed in tumors from female patients with adenocarcinomas without a history of smoking and with Asian ethnicity, but can occur in patients with prior smoking history and across all races and genders (Lynch et al. 2004). The evidence base for the treatments used for the EGFR+ population is more modest than the evidence base for treatments used for EGFR negative NSCLC, which makes development of a model reflective of the latest evidence base more manageable. Future activities may include expanding the model to other subpopulations of interest.

# 3 Purpose

The aim was to develop a flexible open-source simulation model that can be used to estimate the value of alternative sequential treatment strategies for patients with metastatic EGFR+ NSCLC. The IVI-NSCLC(egfr+) model is accessible to both technical and non-technical end-users and allows them evaluating the impact of uncertainty in clinical evidence, alternative model structures, the decision framework of choice (i.e. cost-effectiveness analysis (CEA)) or multi-criteria decision analysis (MCDA)), the inclusion of novel concepts of values (i.e., value of hope), and perspective (healthcare or limited societal) on estimates of value.

#### 3.1 Value assessment

The IVI-NSCLC(egfr+) model is designed to assess the value of multiple competing sequential treatment strategies for patients with metastatic EGFR+ NSCLC starting with 1st line treatment (1L), followed by 2nd line treatment (2L), and treatment beyond 2nd line (2L+) as outlined in Section 7.

The model is suitable for informing decisions for specific (sub)populations, but is not suitable for making predictions at the individual level, nor should it replace the patient-physician shared decision-making process. Local decision makers can modify the model to perform analyses based on populations or parameter values reflective of their local setting while accounting for scientific uncertainty that helps them understand the confidence with which they make decisions.

The IVI-NSCLC(egfr+) model is not a value assessment framework, but a model that simulates the health outcomes, risks, and costs associated with sequential treatment sequences for metastatic EGFR+ NSCLC. It can therefore be used with any value framework preferred by the user. Currently, two methodologies for decision analysis are supported by the model: CEA based on cost per quality adjusted life year (QALY) expressed as net-monetary benefit (NMB) and MCDA (Keeney and Raiffa 1993). The MCDA is implemented according to the methodology described in Thokala et al. (2016) by weighting performance on different criteria (see Section 9.2). As described in as described in the documentation to the IVI-RA model, a linear partial value function is used to translate performance on each criteria to a common scale and an additive model aggregates results. The assessment of value can be performed from a health care sector perspective by only incorporating health care sector costs, or from a (limited) societal perspective by also including productivity costs.

Garrison et al. (2017) suggest five concepts of value that researchers should consider adding to the standard cost per QALY based CEA: (i) a reduction in uncertainty from a diagnostic test; (ii) insurance value for healthy individuals due to reduction against physical risk; (iii) the value of hope for individuals who become risk-loving and would rather pay for a therapy with a long right survival tail than a therapy with a shorter right survival tail but an equivalent (or shorter) expected life-expectancy; (iv) real option value when a therapy allows an individual to benefit from future medical innovations; and (v) scientific spillovers when the benefits of an innovation cannot be entirely appropriated by the innovator.

The value of hope is most relevant for innovations that increase longevity and might be particularly well suited to the analyses of treatments for NSCLC. Traditionally, CEA focuses on maximizing expected QALYs; however, patients might be willing to take gambles (i.e., they become "risk lovers") and care about the variation in benefits and costs, not simply the means. If patients value hope, they may prefer the treatment with greater variability in survival over the treatment with less variability despite having the same expected survival. In contrast, if patients are risk-averse, they

may prefer the latter to avoid an unlucky outcome. Either way, they many have a preference for one intervention over its alternative even though they appear identical based on its average costs and benefits. Patients may place substantial value on a modest chance of a durable survival response, over and above average survival, and decision-makers acting on their behalf may want to consider this aspect when making population level decisions regarding the value of interventions.

The IVI-NSCLC(egfr+) model allows users to incorporate value of hope into their analyses, while noting that the approach is less well-established than conventional cost-effectiveness analysis. Mathematical details are provided in Appendix H. Future versions of the IVI-NSCLC(egfr+) may consider incorporating other value components, such as insurance value.

### 3.2 Evaluation of scientific uncertainty

Decision models can be used to inform efficient use of health care resources, but often lead to scientific disagreements and mistrust among stakeholders. Although (a subset) of model inputs are typically informed by a formal evidence synthesis to ensure all relevant evidence is considered, decisions regarding the mathematical structure relating model inputs to outputs are frequently made without evaluating the impact on findings. While robustness can be assessed by means of sensitivity analyses, these are typically limited to studying the impacts of varying model inputs. For any given disease, a variety of modeling approaches have typically been proposed in the literature. In order to evaluate the impact of these different approaches on estimates of value in a systematic way, flexible open-source models are needed to not only capture the uncertainty in model input parameters (i.e. parameter uncertainty), but also capture alternative model structures (i.e. structural uncertainty). This flexibility facilitates demonstrating the implications of different areas of uncertainty and leads to a better understanding of the reasons why value estimates can vary. Additional details on our approach to uncertainty analysis are provided in Section 11.

# 4 Iterative process

In order for a decision model to remain relevant over time it needs to evolve along with the clinical evidence and scientific insights. An open-source approach facilitates iterative development and collaboration between multiple clinical and methodological experts. We will use a 4-step process for the development of flexible decision or simulation models for value assessment:

- 1. Public release of the model. The initial release of an OSVP model must be flexible and allow users to choose from a large number of plausible model structures and approaches based on clinical practice and previous modeling efforts.
- 2. Invite feedback and suggested improvements to the model in a public comment period.

- 3. A panel of experts determines which of the evidence-based suggestions for improvement suggested in Step 2 should be implemented by means of a modified Delphi process.
- 4. Revise the model based on the feedback from the technical expert panel in Step 3. The 4-step process is designed to be repeated many times so that the modeling approach and evidence considered can be refined over time.

# 5 Components

Version 1 of the IVI-NSCLC(egfr+) model is designed to provide a starting point for open debate. To facilitate transparency, understanding, debate and collaboration among diverse stakeholders, the model consists of the following components available in the public domain:

**Source code:** R code for the model is available in our IVI GitHub repository. Modelers and programmers may adapt the source code for their own purposes or collaborate to improve the code.

R package: The model is released as an R package with documentation available online. Researchers can use the package to run the model for custom analyses. Use of the package is recommended when performing analyses for scientific research.

**Detailed model documentation:** This document provides extensive technical details on the model structure, statistical methods for parameter estimation, and source data.

**Expert Model Interface:** For users not well-versed in the R programming language, a web application for running the model online is provided. The web application is designed for custom analyses and allows users full control over the treatments, patient population, model structures, parameter values, and simulation settings.

Value Tool: An important aim of OSVP is to obtain feedback from as many relevant stakeholders as possible. A general audience web-application has been developed, allowing those who are not experts in modeling or health economics, to interact with the model.

# 6 Population

The model simulates outcomes for a population of patients, each with distinct characteristics. Three characteristics can currently be specified: T790M mutation status, age, and gender. The proportion of patients with a T790M mutation is based on Ma et al. (2011), who report that 82 of 158 (52%) patients tested positive for a mutation after TKI failure. The age distribution is based on the incidence of lung and bronchus cancer cases between 2011 - 2015 as reported by the Surveillance, Epidemiology, and End Results (SEER) Program, which is well approximated by a normal distribution with mean 70.39 and standard deviation 11.68 (Figure A1) (Noone et al.). The same SEER

review reports that women account for 123,650 (48%) of 257,301 incident cases between 2011 and 2015.

Within the model, the proportion of patients with T790M mutations affects the 2L treatments that patients receive. Age and gender impact productivity costs since wages and employment status vary by gender. However, neither age, gender, or any other prognostic factors or effect modifiers impact transition rates or adverse events because these parameters are estimated from randomized controlled trial (RCT) summary data (see Section 10) and not from individual-level data.

## 7 Treatment strategies

Sequential treatment strategies that can be evaluated with version 1 of the model were informed by the available evidence base, guidelines, and clinician input. They are shown in Figure 1.



Figure 1: Sequential treatment strategies of interest to be compared with the model

Tyrosine kinase inhibitors (TKIs) can be used at 1L. If osimertinib is not selected as the 1L treatment, then possible second line treatments (2L) and treatments beyond second line (2L+) depend

on whether a patient acquired a T790M mutation. For example, after discontinuation of erlotinib, patients with a T790M mutation are treated with osimertinib while patients without a mutation can either be treated with platinum-based doublet chemotherapy (PBDC) or a combination of PBDC and anti-VEGF therapy. At 2L+, patients can be treated with a number of combination therapies including those that include immune checkpoint inhibitors (ICIs). Patients cannot be treated with the same treatment at multiple lines.

The specific treatments included in the model are:

• TKIs: erlotinib, gefitinib, afatinib, dacomitinib, osimertinib

• anti-VEGF therapy: bevacizumab

• ICIs: nivolumab, pembrolizumab, atezolizumab

### 8 Model structure

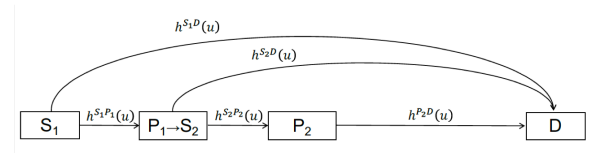
#### 8.1 Disease model

The IVI-NSCLC is an individual-level continuous-time state transition model (CTSTM) in which patients progress through multiple health states. Sequential treatment is modeled using the CTSTM by expanding the number of health states according to the number of treatment lines. In general, we define a health state for each treatment line, a health state after progression on the final line, and a death state, so a model with n treatment lines will have n + 2 health states. A patient with stable disease moves to the second treatment in a sequence after progression on the first treatment, to the third treatment after progression on the second treatment, and so on. A patient can die at any time.

Unfortunately, the evidence base is currently too limited to explicitly model the transition rates by line of treatment and a simplified model structure is needed. Two options are available with the IVI-NSCLC model: a 4-state model (Figure 2) and a three-state model (Figure 3). The 3-state model is consistent with common methods for health economic modeling in oncology. In contrast, the 4-state model incorporates sequential treatment by explicitly modeling the effect of 2L treatment.

In the 4-state model, patients begin 1L treatment in stable disease (S1) and can either transition to the progression state (P1) or death (D). At progression, patients move to the progression-free (stable) state (S2) with 2L treatment and can again either have their disease progress (P3) or die. At P3, patients begin 2L+ treatment and remain in the progressed state until death. Time-varying hazard rates for transitions between states r and s and are denoted by  $h^{rs}(u)$ .

In the three-state model, all hazard rates are based on 1L clinical trial evidence. Patients begin in stable disease (S1) and they can transition to the progression state (P1) or death (D).



 $S_1$ = Progression-free (stable disease) with 1L treatment

 $P_1$  = Progression with 1L treatment

 $S_2$ = Progression-free (stable disease) with 2L treatment

 $P_2$ = Progression with 2L treatment, captures the survival with 2L+ without making a distinction between a progression free and progression phase

D = Dead

 $h^{S_1P_1}(u)$  hazard for transitioning from progression-free to progression with 1L treatment at time u

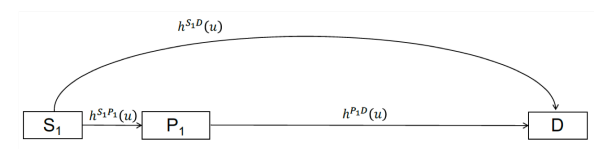
 $h^{S_1D}(u)$ = hazard for transitioning from progression-free to dead with 1L treatment at time u

 $h^{S_2P_2}(u)$ = hazard for transitioning from progression-free to progression with 2L treatment at time u

 $h^{S_2D}(u)$  = hazard for transitioning from progression-free to dead with 2L treatment at time u

 $h^{P_2D}(u)$  = hazard for transitioning from progression on 2L to dead at time u

Figure 2: Model structure with 4 states describing development of disease over time for a sequence starting with 1L, followed by 2L and 2L+ treatment; 2L+ treatment is captured with the L2 progression state



 $S_1$ = Progression-free (stable disease) with 1L treatment

 $P_1$ = Progression with 1L treatment, captures the survival with 2L and 2L+ without making a distinction between progression free and progression phases

D= Dead

 $h^{S_1P_1}(u)$ = hazard for transitioning from progression-free to progression with 1L treatment at time u

 $h^{S_1D}(u)$ = hazard for transitioning from progression-free to dead with 1L treatment at time u

 $h^{P_1D}(u)$ = hazard for transitioning from progression on 1L to dead at time u

Figure 3: Alternative model structure with 3 states describing development of disease over time for a treatment sequence starting with 1L until death; 2L and 2L+ treatment is captured with the progression state

The hazard rate,  $h^{rs}(u)$ , follows different parametric distributions as estimated with a multi-state network meta-analysis (NMA) for non-reversible 3-state models (Section 10.1). As such, the disease model is perfectly integrated with the parameter estimation and the simulation techniques depend on the statistical approach. In general, there are two type of multi-state models: "clock forward" (i.e., Markov) models and "clock reset" (i.e., semi-Markov) models. In the "clock reset" approach, time u in  $h^{rs}(u)$  resets to 0 after each transition whereas in the "clock forward" approach, time u refers to time since the start of the model. If a "clock forward" approach is used, then health state probabilities can be calculated analytically using the Aalen and Johansen (1978) estimator; conversely, if a "clock reset" approach is taken, then individual-level simulation-based approaches must typically be used to estimate health state probabilities (Putter et al. 2007; de Wreede et al. 2011; Jackson 2016).

When a 3-state economic model is used, the 1L NMA is used to directly simulate disease progression and a "clock forward" approach is sufficient. However, if the number of health states is greater than 3, then separate "clock forward" NMAs are fit to estimate line-specific transition rates. For example, in the 4-state model, the 1L results are used to simulate the transition from  $S_1$  to  $P_1 \rightarrow S_2$  and  $S_1$  to  $P_1$ , and the 2L results are used to simulate the transition from  $P_1 \rightarrow P_2$  to  $P_2$ ,  $P_2$  to  $P_3$ , and  $P_1 \rightarrow P_3$  to  $P_3$ . In other words, in the 4-state model, a "clock forward" model is used for 1L but time  $P_1 \rightarrow P_3$  to  $P_3 \rightarrow P_4$  to  $P_4 \rightarrow P_5$  to  $P_5 \rightarrow P_5$  to P

Our statistical approach for modeling transitions between health states also allows us to incorporate potentially relevant prognostic factors or effect-modifiers, although we currently lack the individual-level data needed to do this. That is, we model the parameters of the survival distributions as functions of covariates so that they can vary across individuals and treatments. A given parameter,  $a_{jt}$ , for individual j using treatment t is modeled as  $\alpha_{jt} = g^{-1}(X_{jt}\beta)$  where  $g^{-1}$  (.) is an inverse transformation function. When data becomes available, this approach has a number of potential benefits including estimating health outcomes that are more precisely tailored to a subpopulation and including carry-over effects from one line of treatment to the next.

#### 8.2 Adverse events

The model simulates the probability of a number of distinct adverse events based on a NMA of RCTs (see Section 10.2). Since the trials typically only report the number of proportion of patients experiencing an adverse event we were unable to model adverse events as a function of time on treatment and instead assume that adverse events occur during the first month of treatment as a function of the 1L treatment.

### 8.3 Utility and health care sector costs

Utility depends on the health state and the probability of an adverse event. Adverse events occur during the first month and cause utility decrements, which vary across treatments. After the first month, utility is only a function of the health state.

Health care sector costs include treatment costs (i.e., drug acquisition and administration costs), costs due to inpatient hospitalizations, outpatient costs, and adverse event costs. Inpatient and outpatient costs vary by health state and accrue until death. Adverse event costs—which accrue during the first month—are a weighted sum of the probability of each adverse event and the event's expected cost. Treatment costs at a given line in the sequence accrue until until disease progression or, if dosage depends on time, the duration of the treatment.

### 8.4 Productivity

Productivity costs are currently estimated using the human capital approach (HCA), which measures lost productivity using work time lost. The approach is rooted in economic theory as time away from work is valued at the market wage, which, in a competitive market, reflects the societal value of work. The HCA contrasts with the friction cost approach (FCA), which only measures productivity costs during the time (i.e., the friction period) required to replace workers leaving employment due to illness. A practical limitation of the FCA is that it is difficult to reliably estimate the friction period, which depends on the unemployment rate and consequently varies considerably across individuals and over time. In general, the HCA tends to overestimate productivity costs while the FCA may underestimate them.

Similar to Hanly et al. (2012), time lost from work in the model stems from three sources: temporary disability, permanent disability, and premature death. At cancer onset, patients miss days from work due to temporary disability which reflects work absence due to sick leave. Following sick leave, permanent survivors can then miss additional time from work due to permanent disability. Time lost due to premature death is the difference between the simulated age of death and the retirement age (age 65). Productivity costs are computed by valuing time lost from work at the market wage.

### 9 Model outcomes

The model simulates the health outcomes, adverse events, and costs associated with treatment strategies based on Figure 1, which are combined for value assessment. The model's time horizon can be selected by the user and defaults to a lifetime horizon. The model outcomes are listed in Table 1.

Table 1: Model outcomes

Category	Outcomes
Health outcomes	Health state probabilities; progression free survival & overall survival; quality-adjusted life-years
Adverse events	Diarrhea, dry skin, elevated alanine transaminase, elevated aspartate transaminase, eye problems, paronychia, pneumonitis, pruritus, rash, stomatitis)
Health care sector costs	Drug acquisition and administration costs; adverse event costs; costs from inpatient hospital stays; costs from hospital outpatient or doctor office visits
Non-health care sector costs	Productivity costs
Value assessment	Cost-effectiveness analysis; multi-criteria decision analysis

### 9.1 Health outcomes, adverse events, and costs

The disease model simulates the times that patients transition to and from health states, which, in turn, is used to estimate the probability that a patient is in a given health state at a given point in time following treatment initiation. Progression free survival (PFS) with 1L treatment is defined as time in state  $S_1$  while PFS with 2L is defined as time in state  $P_1 \to S_2$ . Overall survival is time until death. Adverse events are summarized using the probability of occurrence.

Costs are computed for multiple categories and separated into health care sector and non-heath care sector costs as recommended by Sanders et al. (2016). Health care sector costs include treatment costs (i.e., drug acquisition and administration costs), adverse event costs, the costs of inpatient hospital stays, and costs from hospital outpatient and doctor office visits. Non-health care sector costs include productivity costs.

Discounted costs and QALYs over the model's time horizon are computed using the continuous time present value given a flow of state values (i.e., utility and annualized costs), which change as patients transition between health states or as costs vary as a function of time since treatment initiation. Denote the set of M state values for a given patient in the simulation sorted by time as  $z_1, \ldots, z_M$ . Discounted costs and QALYs are then given by,

$$\sum_{m=1}^{M} \int_{u_m}^{u_m+1} z_m e^{-ru} du = \sum_{m=1}^{M} z_m \left( \frac{e^{-ru_m} - e^{-ru_{m+1}}}{r} \right), \tag{1}$$

where r > 0 is the discount rate and u is time. If r = 0, then the present value simplifies to  $\sum_{m=1}^{M} z_m (u_{m+1} - u_m)$ .

#### 9.2 Value assessment

If CEA is used for value assessment, then the value of treatment is estimated using the NMB defined as discounted QALYs multiplied by willingness to pay per QALY less costs. Costs in analyses conducted from a limited societal perspective include both productivity costs and health care sector costs while costs in analyses conducted from a health care sector perspective only include health care sector costs.

Any combination of simulated model outcomes can be used for MCDA. If a 3-state model is selected, the MCDA in IVI's web applications is currently based on the following criteria: progression free survival (PFS) with 1L treatment, post-progression survival, total health care sector costs, oral administration (vs. intravenous administration), and years since FDA approval. If a 4-state model is selected, then PFS with 2L treatment is added as an additional criteria. Similarly, if the MCDA is conducted from limited societal perspective, then productivity costs are added as a criteria. Performance on the oral administration criteria is computed as the percentage of simulated life-years treated using oral therapies rather than intravenous therapies. Performance on the years since FDA approval criteria is the weighted time since FDA approval for each treatment in a treatment sequence, with weights equal to the number of life-years spent using a particular treatment with the sequence. In the web interfaces users can input their own weights for each of the criteria, but it is important to note that we have not conducted the surveys required to elicit weights for a representative sample of patients.

# 10 Source data and parameter estimation

Key parameters for the model relate to: (i) treatment effects in terms of transitions between the health states; (ii) adverse events; (iii) utilities; (iv) healthcare resource use; and (v) productivity. Parameter estimates for the current version of the IVI-NSCLC(egfr+) model are based on currently available published evidence identified by means of a systematic literature review and synthesized with meta-analysis techniques where appropriate. All costs are in 2018 U.S. dollars.

#### 10.1 Treatment effects for transition rates

Relevant evidence to estimate treatment effects with the alternative TKIs and chemotherapy regimens in terms of transitions between the health states was obtained with a systematic literature review of RCTs evaluating the efficacy of relevant competing interventions for the treatment of adult patients with metastatic EGFR+ non-squamous NSCLC by line of treatment (i.e. 1L, 2L, and 2L+) in terms of OS and PFS. Details regarding eligibility criteria, search strategies, included

studies, extracted information regarding study and patient characteristics, results, and KM curves are provided in (Appendix X). The identified evidence base was used to 1) estimate the relative treatment effects for each 1L treatment versus a defined reference treatment; 2) estimate the absolute effects with the corresponding 1L reference treatment; and 3) estimate the absolute treatment effects with 2L treatment options.

#### 10.1.1 Relative treatment effects with 1L therapy

Relative treatment effects of the competing TKIs in comparison to gefitinib for the 1L treatment of EGFR+ NSCLC were estimated by means of a Bayesian network meta-analysis (NMA) using alternative fixed and random effects models.

RCTs are considered the gold standard to assess treatment effects of medical interventions. For many disease states there are multiple competing interventions to choose from. However, an RCT will rarely include all competing interventions of interest for a particular disease state when there are more than two relevant treatment options available. As such, an individual trial rarely provides all the information needed to guide evidence-based treatment selection. When each RCT compares only a subset of the interventions of interest—it may be possible to represent the evidence base as a network where all trials have at least one intervention in common with another trial. Such a network of trials involving treatments compared directly, indirectly, or both, can be synthesized by means of a network meta-analysis (add reference). As a result we not only get pooled results of available treatment comparisons studied in a head-to-head fashion but also relative treatment effects between interventions not compared directly. In order to ensure that a network meta-analysis are not affected by bias due to differences in prognostic factors between studies, we want to only consider the relative treatment effects of each trial. This implies that all interventions have to be part of one network of trials where each trial has at least one intervention in common with another trial.

Patients are randomized only within trials, not across trials, so there is a risk that patients participating in different trials differ with respect to demographic, disease or other characteristics. In addition, features of the trials themselves may differ. If these trial or patient characteristics are effect modifiers, i.e. they affect the treatment effects of an intervention versus a control, then there are systematic differences in treatment effects across trials. Systematic differences in known and unknown effect-modifiers among studies comparing the same interventions in direct fashion result in between-study heterogeneity. An imbalance in the distribution of effect modifiers between studies comparing different interventions will result in transitivity or consistency violations and therefore biased indirect comparisons.

The RCTs that were part of one evidence network and were deemed sufficiently similar were included in the NMA. The evidence network for 1L treatments is presented in Figure 4. Information regarding

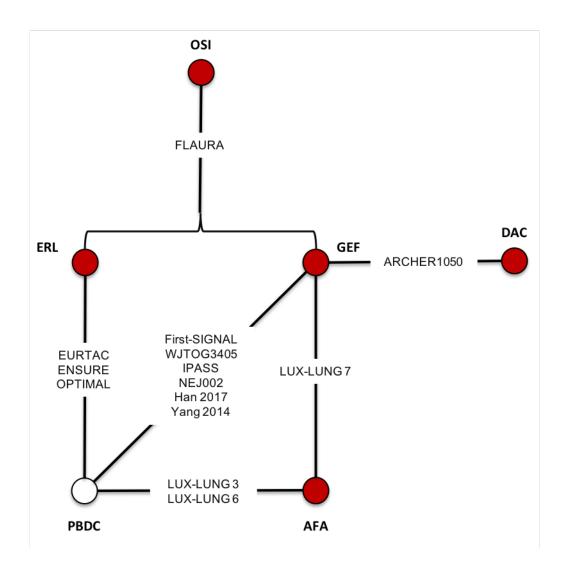


Figure 4: Network of RCTs to estimate relative treatment effects regarding PFS and OS with 1L therapies

the distribution of patients characteristics across the studies in the network are presented in (Appendix X).

### 10.1.1.1 Network meta-analysis model

Traditionally, NMAs in oncology are based on separate analyses for OS and PFS, with relative treatment effects estimated using reported hazard ratios (HR). There are two limitations of this approach. First, HR estimates rely on the proportional hazard assumption, which is implausible if the hazard functions of the competing interventions cross (Dias et al. 2018). Second, separate analyses of OS and PFS ignore the structural relationship between stable disease, progression, and death and implicitly assume that transition probabilities for stable to death and progression to

death are equal.

We overcome these limitations by introducing a multi-state NMA that explicitly estimates each possible transition in a 3-state model—stable to progression, stable to death, and progression to death—and modeling time-varying hazard rates and relative treatment effects with known parametric survival functions or fractional polynomials. Our approach is illustrated in Figure 5 and expressed mathematically as follows:

$$\ln \left( h_{ik}^{SP}(u) \right) = \begin{cases} \alpha_{1ik} + \alpha_{2ik} u^{p_1} + \alpha_{3ik} u^{p_2} & \text{if } p_1 \neq p_2 \\ \alpha_{1ik} + \alpha_{2ik} u^p + \alpha_{3ik} u^p \ln(u) & \text{if } p_1 = p_2 = p \end{cases}$$

$$\ln \left( h_{ik}^{SD}(u) \right) = \alpha_{4ik}$$

$$\ln \left( h_{ik}^{PD}(u) \right) = \alpha_{5ik} + \alpha_{6ik} u^{p_1}$$

$$\begin{pmatrix} \alpha_{1ik} \\ \alpha_{2ik} \\ \alpha_{3ik} \\ \alpha_{4ik} \\ \alpha_{5ik} \\ \alpha_{6ik} \end{pmatrix} = \begin{pmatrix} \mu_{1ik} \\ \mu_{2ik} \\ \mu_{3ik} \\ \mu_{4ik} \\ \mu_{5ik} \\ \mu_{6ik} \end{pmatrix} + \begin{pmatrix} \delta_{1,ik} \\ \delta_{2,t_{ik}} - d_{2,t_{i1}} \\ 0 \\ 0 \\ d_{3,t_{ik}} - d_{3,t_{i1}} \\ 0 \end{pmatrix}$$

$$\delta_{1,ik} \sim N(d_{1,t_{ik}} - d_{1,t_{i1}}, \sigma_{d_1}^2)$$

$$(2)$$

where 
$$u^0 = \ln(u)$$
 and  $d_{1,1} = 0$ ,  $d_{2,1} = 0$ , and  $d_{3,1} = 0$ 

 $h_{ik}^{SD}(u), h_{ik}^{PD}(u)$ , and  $h_{ik}^{SD}(u)$  are the the hazard rate for disease progression, dying post-progression, and dying pre-progression, respectively, in study i for treatment arm k at time u. The  $\alpha_{\cdot ik}$  are regression coefficients that represent the scale and shape parameters of the log hazard function in study i for treatment arm k. The  $\mu_{\cdot i}$  reflect the study effects regarding the scale and shape parameters in each study i. The  $\delta_{1,ik}$  are the study specific true underlying relative treatment effects regarding the scale of the log hazard function for the transition from stable to progression, which are described by a normal distribution with an average effect for treatment t relative to reference treatment 1,  $d_{1,1t}$ , and between study heterogeneity  $\sigma_{d_1}^2$ .  $d_{2,1t}$  are the treatment specific relative effects regarding the first shape parameter of the log hazard function for the transition from stable to progression relative to the reference treatment 1 (i.e. gefitinib). To facilitate parameter estimation, we assumed that treatment has no effect on the 2nd shape parameter. Transition rates between stable disease and death are likely to be relatively small and therefore assumed to be independent of time and the same for all treatments.  $d_{3,1t}$  are the treatment specific relative effects regarding the scale parameter of the log-hazard function for the transition from progression to death

$$S_{ik}(u) \xrightarrow{h_{ik}^{SP}(u)} P_{ik}(u) \xrightarrow{h_{ik}^{PD}(u)} D_{ik}(u)$$

 $S_{ik}(u)$  = progression -free (stable disease) in study i, treatment arm k at time u

 $P_{ik}(u)$ = progressed disease in study i, treatment arm k at time u

 $D_{ik}(u)$ = dead in study i, in treatment arm k at time u

 $h_{ik}^{SP}(u)$ = hazard rate for disease progression in study i, in treatment arm k at time u

 $h_{ik}^{PD}(u)$ = hazard rate for dying post-progression in study i, in treatment arm k at time u

 $h_{ik}^{\mathit{SD}}(u)$ = hazard rate for dying pre-progression in study i, in treatment arm k at time u

Figure 5: Relationship between stable disease (S), progression (P) and death (D) as used in the multi-state network meta-analysis model

relative to reference treatment 1 (i.e. gefitinib).

When  $p_1 = 0$  and  $\alpha_{3ik} = 0$  the log-hazard functions for the transitions between stable disease and progression, and progression and death follow a Weibull distribution. When  $p_1 = 1$  and  $\alpha_{3ik} = 0$  these log-hazard functions follow a Gompertz distribution. When  $\{(p_1, p_2) = (0, 0), (0, 1)\}$  and  $\alpha_{3ik} \neq 0$ , the log-hazard functions follow a second order fractional polynomial that are extensions of the Weibull and Gompertz model to allow for arc- and bathtub shaped log-hazard functions.

A fixed effects model is obtained by replacing  $\delta_{1,ik} \sim N(d_{1,t_{ik}} - d_{1,t_{i1}}, \sigma^2)$  with  $\delta_{1,ik} = d_{1,t_{ik}} - d_{1,t_{i1}}$ .

If it is assumed that treatment has only a direct effect on the transitions from stable to progression, which can be reasonably defended when a particular treatment upon disease progression is discontinued, the model can be simplified by setting  $d_{3,t_{i1}} = 0$ .

### 10.1.1.2 Likelihood

The model parameters were estimated based on the number of patients r in each of the three health states (stable, progression and death) at time u for each arm k of each trial i obtained from the published Kaplan-Meier (KM) curves for time points for which both PFS and OS were reported. Accordingly, a multinomial likelihood was used for the proportion of patients in each of the three health states at any time point for each arm of each trial  $(S_{ik}(u), P_{ik}(u), \text{ and } D_{ik}(u))$  according to:

$$(r_{ik}^S(u), r_{ik}^P(u), r_{ik}^D(u)) \sim multinomial(S_{ik}(u), P_{ik}(u), D_{ik}(u), n_{ik}(u))$$

$$(3)$$

where 
$$n_{ik}(u) = r_{ik}^{S}(u) + r_{ik}^{P}(u) + r_{ik}^{D}(u)$$
 and  $S_{ik}(u) + P_{ik}(u) + D_{ik}(u) = 1$ 

Arbitrary hazard rate functions can be approximated with a set of discontinuous constant hazard rates over successive time intervals. The total follow-up time can be partitioned into M successive non-overlapping intervals indexed by m = 1, ..., M. We refer to interval m as  $U_m$  and write  $u \in U_m$  to denote  $u_m \le u < u_{m+1}$ . The length of  $U_m$  is  $\Delta u_m = u_{m+1} - u_m$ .

For each interval m, the proportions  $S_{ik}(u)$ ,  $P_{ik}(u)$ , and  $D_{ik}(u)$  are related to the time-varying hazards  $h_{ik}^{SP}(u)$ ,  $h_{ik}^{SD}(u)$ ,  $h_{ik}^{PD}(u)$  according to the following set of differential equations (Jansen and Trikalinos 2013):

$$S_{ik}(u) = S_{ik}(u_m)e^{-(h_{ikm}^{SP} + h_{ikm}^{SD})(u - u_m)}$$

$$P_{ik}(u) = P_{ik}(u_m)e^{-h_{ikm}^{PD}(u - u_m)} + \frac{S(u_m)h_{ikm}^{SP}(e^{-(h_{ikm}^{SP} + h_{ikm}^{SD})(u - u_m)} - e^{-h_{ikm}^{PD}(u - u_m)})}{h_{ikm}^{PD} - h_{ik}^{SP} - h_{ikm}^{SD}}$$

$$D_{ik}(u) = 1 - S_{ik}(u) - P_{ik}(u)$$

$$(4)$$

In order to estimate  $h_{ikm}^{SP}$ ,  $h_{ikm}^{SD}$ , and  $h_{ikm}^{PD}$  for each interval m, data at three time points were used:  $u_m$ ,  $u_{m+\frac{1}{2}}$ , and  $u_{m+1}$ . The obtained estimates of the hazards for interval m were assigned to the time point corresponding to the mid point of the interval when used in the NMA.

For time points of a trial for which only OS data was reported (and no PFS), the number of patients that survived time interval m ( $r_{ikm}^{OS}$ ) out of the number of patients alive at the beginning of interval ( $n_{ikm}^{OS}$ ) were used for the model parameter estimation. A binomial likelihood was used for the conditional survival probabilities ( $p_{ikm}^{OS}$ ):

$$r_{ikm}^{OS} \sim binomial(p_{ikm}^{OS}, n_{ikm}^{OS}) \tag{5}$$

The conditional survival probabilities inform  $h_{ik}^{SD}(u)$  and  $h_{ik}^{PD}(u)$  according to:

$$p_{ikm}^{OS} = e^{-(h_{ikm}^{SD} + h_{ikm}^{PD})(u - u_m)}$$
(6)

where the obtained estimates of the hazards for interval m assigned to the time point corresponding

to the mid point of the interval when used in the NMA.

#### 10.1.1.3 Prior distributions

The following prior distributions for the parameters of the model expressed with Equation 2 were used:

$$\begin{pmatrix} \mu_{1i} \\ \mu_{2i} \\ \mu_{3i} \\ \mu_{4i} \\ \mu_{5i} \\ \mu_{6i} \end{pmatrix} \sim MVN \begin{pmatrix} \begin{pmatrix} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \end{pmatrix}, T_{\mu} \end{pmatrix} \qquad T_{\mu} = \begin{pmatrix} 100 & 0 & 0 & 0 & 0 & 0 \\ 0 & 10 & 0 & 0 & 0 & 0 \\ 0 & 0 & 10 & 0 & 0 & 0 \\ 0 & 0 & 0 & 100 & 0 & 0 \\ 0 & 0 & 0 & 0 & 100 & 0 & 0 \\ 0 & 0 & 0 & 0 & 0 & 10 & 0 \\ 0 & 0 & 0 & 0 & 0 & 0 & 10 \end{pmatrix}$$

$$\begin{pmatrix} d_{1,1t} \\ d_{2,1t} \\ d_{3,1t} \end{pmatrix} \sim MVN \begin{pmatrix} \begin{pmatrix} 0 \\ 0 \\ 0 \\ 0 \end{pmatrix}, T_{d} \end{pmatrix} \qquad T_{d} = \begin{pmatrix} 2 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 2 \end{pmatrix}$$

$$\sigma_{d_{1}} = uniform(0.2)$$

$$(7)$$

The prior distribution for the probabilities at the beginning of each interval,  $u_m$  was:

$$(S_{ik}(u_m), P_{ik}(u_m), D_{ik}(u_m)) \sim Dirichlet(0, 0, 0)$$
 (8)

#### 10.1.1.4 Estimated time-varying hazard ratios

To be added

Figure 6: First line estimates of hazard ratios from stable to progression relative to gefitinib from the multi-state network meta-analysis

#### 10.1.2 Absolute effects with 1L reference treatment

Absolute effects with 1L therapy with the reference intervention, i.e. gefitinib, were estimated with the multi-state meta-analysis model expressed in Equation 9 with likelihoods and transformations

according to Equation 3, Equation 4, Equation 5, and Equation 6 based on the reported KM curves of all the gefitinib treatment arms of the available RCTs.

$$\ln \left( h_i^{SP}(u) \right) = \begin{cases} M_1 + M_2 u^{p_1} + M_3 u^{p_2} & \text{if } p_1 \neq p_2 \\ M_1 + M_2 u^p + M_3 u^p \ln(u) & \text{if } p_1 = p_2 = p \end{cases}$$

$$\ln \left( h_i^{SD}(u) \right) = M_4$$

$$\ln \left( h_i^{PD}(u) \right) = M_5 + M_6 u^{p_1}$$

$$(9)$$

where  $u^0 = \ln(u)$ 

 $h_i^{SD}(u)$ ,  $h_i^{PD}(u)$ , and  $h_i^{SD}(u)$  are the underlying hazard rates for disease progression, dying postprogression, and dying pre-progression, respectively, in study i at time u. The M represent the pooled scale and shape parameters describing the log hazard functions over time. As before, when  $p_1 = 0$  and  $M_3 = 0$  the log-hazard functions for the transitions between stable disease and progression, and progression and death follow a Weibull distribution. When  $p_1 = 1$  and  $M_3 = 0$  these log-hazard functions follow a Gompertz distribution. When  $\{(p_1, p_2) = (0, 0), (0, 1)\}$  and  $M_3 \neq 0$ , the log-hazard functions follow a second order fractional polynomial to allow for arc- and bathtub shaped log-hazard functions.

Figure 7 shows the hazard functions for the transitions between stable, progression, and death with gefitinib for each fitted model.

The posterior distributions of M along with the parameters representing the relative treatment effects of each TKI relative to gefitinib  $(d_{1,t})$ , were used to simulate a posterior distribution for PFS and OS as displayed in Figure 8 for each 1L treatment and each fitted model.

Figure 7: First line estimates of hazard rates over time for transitions between S, P and D with gefitinib from the multi-state meta-analysis



Figure 8: First line estimates of progression-free survival and overall survival for the competing interventions obtained with the multi-state (network) meta-analysis

Notes: The simulated posterior distribution of the parameters of the Bayesian multi-state NMA was used to simulate a distribution of progression-free survival and overall survival curves. The curves in the figure are posterior means. Curves with credible intervals are shown in the appendix.

#### 10.1.3 Absolute effects with 2L therapy

For the 2L treatment options, the available number of RCTs among patients who progressed on 1L TKI were limited. For T790M positive patients, the transition rates with osimertinib were estimated based on data from two RCTs according to Equation 9. Regarding other possible treatments, there was only data available for PBDC from three RCTs, which were used to estimate transition rates according to Equation 9 as well. In the absence of RCT evidence, these estimates were assumed to be representative of the other 2L treatment regimens presented in Figure 1 for both an all-comer population and a T790M mutation negative population. Figure 9 shows the hazard functions for the transitions between stable, progression, and death for each fitted model and Figure 10 shows the corresponding PFS and OS curves.

Figure 9: Second line estimates of hazard rates over time for transitions between S, P and D from the multi-state meta-analysis

#### **10.1.4** Software

The parameters of the different models were estimated using a Markov Chain Monte Carlo (MCMC) method implemented in the JAGS software package. All JAGS analyses were run using R statistical software (R Core Team 2014).

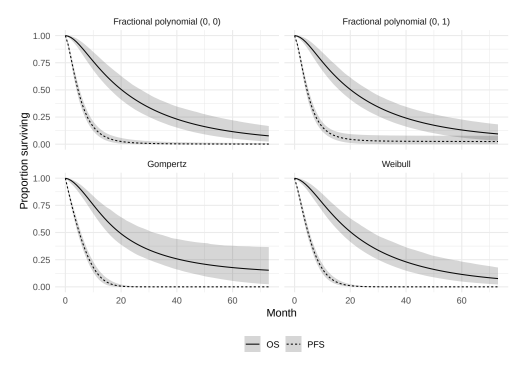
#### 10.1.5 Model selection

The residual deviance and the deviance information criterion (DIC) were used to compare the goodness-of-fit of the competing NMA models, i.e. fixed and random effects Weibull, Gompertz, and fractional polynomial models (See (Appendix X)) (Dias et al. 2018). The DIC provides a measure of model fit that penalizes model complexity. In general, a more complex model results in a better fit to the data, demonstrating a smaller residual deviance. The model with the better trade-off between fit and parsimony has a lower DIC. A difference in the DIC of about 10 points is considered meaningful.

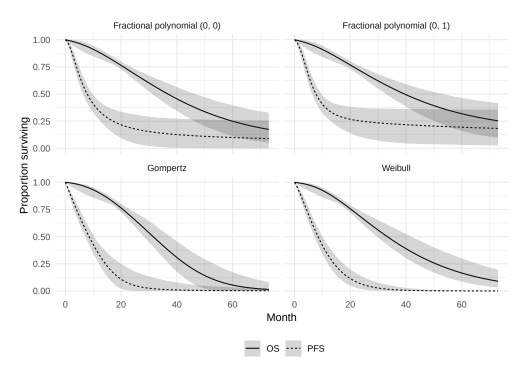
Informed by model fit criteria, we used treatment effect estimates obtained with the fixed and random effects, Weibull and fractional polynomial models without a treatment effect for progression to death for the economic model.

### 10.2 Adverse events

The identified RCTs also provided the evidence base to estimate treatment specific adverse event probabilities. Since adverse events in nearly all RCTs were reported as the number of patients



#### (a) Platinum based doublet chemotherapy



(b) Osimertinib among T790M positive patients

Figure 10: Second line estimates of progression-free survival and overall survival from the multi-state meta-analysis

Notes: The simulated posterior distribution of the parameters of the Bayesian multi-state NMA was used to simulate a distribution of progression-free survival and overall survival curves. The solid lines are posterior means and the shaded region denotes the 95 percent credible interval.  $\frac{1}{28}$ 

experiencing the event, the NMA was performed on the proportion of patients experiencing the event of interest with a binomial likelihood and logit link (Dias et al. 2018, Chapter 2). Add details about pooling and the priors...

#### 10.3 Utilities

Health state utilities based on public literature are reported in Table 2. We assume that utilities do not vary across treatment strategies. We use estimates from Nafees et al. (2017), a global study of patients with metastatic NSCLC that used health states representative of 1L treatment to derive utility values. Estimates are based on the "global" analysis that pooled estimates across 6 countries (Australia, China, France, Korea, Taiwan, and the UK) for patients with stable disease and no side effects. Utilities for patients with progressed disease are based on Nafees et al. (2008), a study representative of metastatic NSCLC patients on 2L treatment. Estimates are from predictions using the mixed model among patients with no adverse events. Utility in the P1/S2 state is estimated using patients that have not yet progressed while on 2L, while estimates in the P2 state are among the progressed 2L patients.

Table 2: Utility by health state

Health state	Mean	Standard error	Reference
S1	0.7540	0.0000	Nafees et al. (2017)
P1/S2	0.6532	0.0222	Nafees et al. (2008)
P2	0.4734	0.0311	Nafees et al. (2008)

Adverse event disutilities are displayed in Table 3. When we cannot identify a disutility for an adverse event, we assume that the disutility is equivalent to the disutility from the "most similar" adverse event. Elevated alanine transaminase and elevated aspartate transaminase are assumed to have no effect on utility since they are based on test results rather than symptoms.

### 10.4 Health care sector costs

#### 10.4.1 Treatment costs

Treatment costs are a function of drug acquisition and drug administration costs. Dosage for each drug based on Federal Drug Administration (FDA) labels is presented in Table 4. Patients using PBDC are, by default, assumed to use a combination of cisplatin and pemetrexed, although this can be adjusted in the R package. At a given treatment line, patients continue to take a drug until disease progression or the end of a treatment cycle. For example, a patient using erlotinib at 1L would take a 150 mg tablet each day until disease progression, at which point they would

Table 3: Disutility due to adverse events

Adverse event	Mean	Standard error	Reference
Elevated alanine transaminase	0.0000	0.0000	N/A
Elevated aspartate transaminase	0.0000	0.0000	N/A
Diarrhea	0.2220	0.0730	Nafees et al. (2017)
Dry skin	0.1510	0.0500	Nafees et al. (2017)
Eye problems	0.1380	0.0460	Nafees et al. (2017)
Paronychia	0.1510	0.0500	Nafees et al. (2017)
Pneumonitis	0.1650	0.0540	Doyle et al. (2008)
Pruritus	0.1510	0.0500	Nafees et al. (2017)
Rash	0.1510	0.0500	Nafees et al. (2017)
Stomatitis	0.1510	0.0500	Nafees et al. (2017)

begin to use the 2L drug. Conversely, a patient using PBDC at 2L would discontinue cisplatin and pemetrexed after completing 6 21-day cycles.

Table 4: Drug dosage

Drug	Dosage
erlotinib	150 mg orally, once daily
gefitinib	250 mg orally, daily
afatinib	40 mg orally, once daily
dacomitinib	45 mg orally, once daily
osimertinib	80 mg orally, once daily
cisplatin	75 mg/m2, $1 x/cycle$ , 6 21-day cycles
pemetrexed	500  mg, 1x/cycle, 6 21-day cycles
bevacizumab	15 mg/kg IV every 3 weeks with carboplatin/paclitaxel
nivolumab	240 mg every 2 weeks or 480 mg every 4 weeks
pembrolizumab	200 mg every 3 weeks
atezolizumab	1200 mg as an IV infusion over 60 min every 3 weeks

In some cases, patients might be required to use multiple dosage forms to obtain the recommended dosage. In these cases, we assume that patients use the cheapest possible combination of dosage forms. For instance, dosage for nivolumab is 240 mg every 2 weeks, so patients are assumed to use 2 100mg vials and 1 40mg vials.

Dosing with cisplatin depends on body weight. The mean body surface area (BSA) in the United States is  $1.6m^2$  for women and  $1.9m^2$  for men, which implies a dose of 120 mg for women of and 142.5 mg for men. The model assumes that there is no vial sharing, so patients use 1 100mg vial and 1 50 mg vial of cisplatin.

Wholesale acquisition costs (WACs) are shown in Table 5. Drug acquisition costs in the model are

equal to the WACs adjusted for discounts and rebates. These discounts can be applied to uniquely to each drug in the R package, but are, by default, assumed to range from 20% to 30%. When historical data was available, we used the most recently available WAC from either ProspectoRx or AnalySource. There was no historical data for dacomitinib, so we used the publicly announced WAC of \$12,400 per month.

Table 5: Wholesale acquisition costs

Drug	Strength	Acquisition cost
erlotinib	150mg tablet	281.71
gefitinib	250mg tablet	257.06
afatinib	40mg tablet	235.05
dacomitinib	45mg tablet	407.39
osimertinib	80mg tablet	487.22
cisplatin	$50 \mathrm{mg}/50 \mathrm{ml}$ vial	14.51
cisplatin	$100 \mathrm{mg} / 100 \mathrm{ml} \ \mathrm{vial}$	35.00
cisplatin	$200 \mathrm{mg}/200 \mathrm{ml}$ vial	100.00
pemetrexed	100mg vial	676.52
pemetrexed	500 mg vial	3382.60
bevacizumab	$100 \mathrm{mg}/4 \mathrm{ml}$ vial	199.24
bevacizumab	$400 \mathrm{mg}/16 \mathrm{ml}$ vial	199.24
nivolumab	$100 \mathrm{mg} / 10 \mathrm{ml} \ \mathrm{vial}$	262.21
nivolumab	$40 \mathrm{mg}/4 \mathrm{ml}$ vial	262.21
pembrolizumab	$100 \mathrm{mg}/4 \mathrm{ml}$ vial	1162.41
atezolizumab	$1200 \mathrm{mg}/20 \mathrm{ml}$ vial	444.03

Table 6 reports drug administration costs. Drug administration costs are based on US Current Procedures Terminology (CPT) codes and accrued each time a patient takes a drug.

Table 6: Drug administration costs

Drug	Administration cost	Source
erlotinib	0.00	N/A
gefitinib	0.00	N/A
afatinib	0.00	N/A
dacomitinib	0.00	N/A
osimertinib	0.00	N/A
cisplatin	91.72	CPT 96417, 96415
pemetrexed	136.15	CPT 96413
bevacizumab	91.72	CPT 96413, 96415, 96417
nivolumab	136.15	CPT 96413
pembrolizumab	136.15	CPT 96413
atezolizumab	136.15	CPT 96413

### 10.4.2 Inpatient costs

Inpatient costs with stable disease are based on the commercial cost estimates from Graham et al. (2018), who separate costs due to adverse events from costs due to adverse events. Monthly costs include those that accrue due to unscheduled hospital stays, intensive care unit visits, and emergency department visits. Inpatient costs with progressed disease are from the estimates reported in Skinner et al. (2018). Costs on 1L treatment are equal to the hospitalization costs during the post-progression period for patients being treated with systemic anti-cancer therapy. Costs on 2L treatment are based on "overall" hospitalization costs during the post-progression period, which are a mixture of costs for those treated with systemic anti-cancer therapy and those untreated.

Table 7: Monthly inpatient medical costs

Health state	Mean	Standard error	Reference
S1	1,909	469	Graham et al. (2018)
P1/S2	5,805	606	Skinner et al. (2018)
P2	7,710	805	Skinner et al. (2018)

### 10.4.3 Outpatient costs

As with inpatient costs, outpatient costs with stable disease are based on Graham et al. (2018) and costs following progression are from Skinner et al. (2018). Outpatient costs estimated by Graham et al. (2018) consist of costs from outpatient visits (e.g., a visit to the general practicioner) and outpatient interventions (e.g. a computed tomography scan). Costs during the post-progression period with 2L treatment are the sum of costs from procedures and physician office visits for those treated with systemic anti-cancer therapy. Costs during the post-progression period with 2L+ treatment are also the sum of costs from procedures and physician office visits, but include patients untreated with systemic anti-cancer therapy in addition to patients treated.

Table 8: Monthly outpatient medical costs

Health state	Mean	Standard error	Reference
S1	63	15	Graham et al. (2018)
P1/S2	1,624	170	Skinner et al. (2018)
P2	1,287	134	Skinner et al. (2018)

#### 10.4.4 Adverse event costs

Costs due to adverse events are taken from a variety of sources. First, we use results from Wong et al. (2018), who estimated the incremental costs of severe (defined as requiring an inpatient stay) adverse

events among patients from cancers of the breast, digestive organs and peritoneum, genitourinary organs, lung, lymphatic and hematopoietic tissue, and skin. The analysis used the Truven Health Analytics Market Scan database to estimate the difference in costs between treatment espidoes involving adverse events and matched treatment episodes without adverse events. Second, Bilir et al. (2016) is used to estimate costs due to test results for elevated alanine transaminase or elevated aspartate transaminase. To remain consistent with our other estimates we only use inpatient costs that accrued during the hospitalization associated with the adverse event, which were based on an analysis of Optum claims data among patients with metastatic melanoma. Finally, Medicare-Severity Diagnosis Related Group (MS-DRG) was used for paronychia, which was not reported in the other studies.

Table 9: Costs of adverse events

Adverse event	Mean	Lower	Upper	Reference
Diarrhea	18,011	15,371	20,499	Wong et al. (2018)
Dry skin	1,025	115	2,011	Wong et al. (2018)
Elevated alanine transaminase	21,411	8,863	33,959	Bilir et al. (2016)
Elevated aspartate transaminase	21,411	8,863	33,959	Bilir et al. (2016)
Eye problems	34,882	23,916	53,943	Wong et al. (2018)
Paronychia	8,249	$6,\!599$	9,898	DRG 602
Pneumonitis	23,922	$22,\!578$	$25,\!554$	Wong et al. (2018)
Pruritus	$28,\!551$	3,996	$67,\!642$	Wong et al. (2018)
Rash	17,137	12,343	23,316	Wong et al. (2018)
Stomatitis	19,801	16,484	24,591	Wong et al. (2018)

### 10.5 Productivity

Temporary disability is measured using the mean duration of absence from work due to sick leave. A central estimate from the literature of the number of missed work days due to cancer is 151 (Mehnert 2011), which we, by default, vary by 20% in either direction in the uncertainty analysis. Permanent disability is measured as the average reduction in work hours among cancer survivors. Short et al. (2008) estimate an average reduction of 3 to 5 hours per week, which is inclusive of survivors who stopped working completely and similar across males and females.

Wages stratified by gender and employment status are reported in Table 10. Estimates of both the weekly wage and the proportion of individuals by employment status are from the U.S. Bureau of Labor Statistics.

#### 10.6 Value of hope

As described in more detail in Appendix H, the value of hope depends on the form of the utility

Table 10: Weekly wages by gender and employment status

Employment status	Percentage	Weekly wage
	Women	
Full-time	72.4%	\$796
Part-time	23.3%	\$326
Unemployed	4.3%	\$0
	Men	
Full-time	84.1%	\$973
Part-time	11.5%	\$321
Unemployed	4.4%	\$0

function and the estimate of relative risk aversion. Following Shafrin et al. (2017), we use the constant relative risk aversion utility function  $u(x) = x^{\eta}$  where  $\eta = 1.39$ , which was estimated among patients with NSCLC.

### 11 Simulation and uncertainty analysis

The individual-level CTSTM is simulated using the general purpose R package **hesim**, which can simulate "clock reset", "clock forward", and mixtures of "clock reset" and "clock forward" models. All economic models in **hesim** are inherently Bayesian, as uncertainty in the parameters from the statistical models are propagated throughout the economic model with probabilistic sensitivity analysis (PSA). Furthermore, since the economic models developed in **hesim** are integrated with the statistical models for parameter estimation, structural uncertainty analysis can be peformed in a straightforward manner by fitting models with different parametric distributions or model structures.

#### 11.1 Individual-level simulation

The individual-level CTSTM simulates trajectories through multiple health states for individual patients. The simulation starts at time 0 in stable disease  $(S_1)$ . Times to health states that can be transitioned to (e.g.,  $P_1 \to S_2$  and D from  $S_1$  in the 4-state model) are then sampled and a patient transitions to the state with the shortest sampled time. The process repeats itself as patients progress though the multi-state model until the time horizon is reached or death. The simulation generates a vector of G simulated health states  $h_1, \ldots, h_G$  and the times that they were entered  $0, \ldots, u_G$  for each patient.

In a "clock forward" model, time does not reset each time a patient enters a new health state. Times to the competing events are consequently sampled from truncated distributions with lower bound equal to the current time u in the simulation. State  $S_1$  at time 0 is an exception, however, as times-to-event are sampled from non-truncated distributions for computational efficiency.

In the mixture of "clock forward" and "clock reset" models, time only resets at certain health states dictated by the multi-state NMA. For example, in the 4-state model, time resets once patients enter state  $S_1 \to P_2$ . In other words, times-to-event are drawn from non-truncated distributions from state  $S_1$  at time 0 and when time resets in state  $S_1 \to P_2$ , but from truncated distributions while in state  $P_2$ . The lower bound of the truncated distribution used from state  $P_2$  is equal to the elapsed time from  $S_1 \to P_2$  to  $P_2$ .

Times-to-event from parametric distributions with known density functions (e.g., Weibull) are fast using standard random number generation algorithms. However, different random number generation approaches are needed for flexibly parametric distributions with density functions that must be computed numerically. For example, the fractional polynomial NMA models the log hazard rate and the cumulative density function (CDF) is computed from the cumulative hazard with numerical integration. There are a number of ways to sample from an arbitrary CDF including the inverse CDF method, sampling from a disrete grid, rejection sampling, and the Metropolis-Hastings algorithm. We current use the discrete grid approach—which we have found to be numerically stable—by sampling time-to-event based on probabilities computed from the CDF.

### 11.2 Parameter uncertainty

Parameter uncertainty is quantified using probabilistic sensitivity analysis (PSA), which propagates uncertainty in the model input parameters throughout the model by randomly sampling the input parameters from suitable probability distributions (Baio and Dawid 2015; Claxton et al. 2005). Probability distributions are determined according to the distributional properties of the statistical estimates, which, in turn, depend on the statistical techniques used, and the distributions of the underlying data. Table 11 displays the probability distribution that we used for the model parameters. When conducting a CEA, the results of the PSA are summarized using standard measures from the literature including cost-effectiveness planes (Black 1990; Barton et al. 2008), cost-effectiveness acceptability curves (CEACs) (Van Hout et al. 1994; Briggs 1999; Fenwick et al. 2001; Barton et al. 2008), the cost-effectiveness acceptability frontier (CEAF) (Barton et al. 2008), and estimates of the expected value of perfect information (EVPI) (Fenwick et al. 2001; Barton et al. 2008). Furthermore, all point estimates from a CEA or MCDA are reported along with 95 percent credible intervals.

### 11.3 Structural uncertainty

Structural uncertainty is assessed in two ways. First, as discussed in Section 10.1 different model specifications were used in the multi-state NMA. These included a Weibull model and two second

Table 11: Probability distributions for probabilistic sensitivity analysis

Parameter(s)	Distribution		
Health state transitions			
Bayesian multi-state NMA	Simulated posterior distribution		
Adverse events			
	C:		
Bayesian NMA for adverse events	Simulated posterior distribution		
Utility			
Health state utility	Normal		
V			
Adverse event disutilities	Normal		
Health care sector costs			
Drug acquisition and administration cost	Fixed		
Discounts and rebates applied to drug acquisition	Uniform		
cost	Ciniorin		
Cost			
Inpatient costs	Gamma		
•			
Outpatient costs	Gamma		
Adverse event costs	Normal		
Productivity costs			
· · · · · · · · · · · · · · · · · · ·	Uniform		
Missed work days	Ulliotiii		
Reduction in hours worked	Uniform		

order fractional polynomial models  $\{(p_0, p_1) = (0, 0), (0, 1)\}$ . A Gompertz distribution was also considered but the model fit was considerably worse.

Second, both 4-state (Figure 2) and 3-state (Figure 3) model structures can be used. The 3-state approach is the typical model structure in economic modeling in oncology and therefore included for consistency with the standard approach. That said, the 4-state model may be preferred because it is able to overcome some of the limitations of the 3-state model. For one, estimates of transition rates following progression are based on specific 2L treatments rather than an "average" of 2L treatments (as in the  $P_1$  to D transition in the 3-state model). Similarly, treatment costs after progression are based on explicit treatment sequences (i.e., 2L treatment following by 2L+ treatment) wheras in the 3-state model assumptions must be made such as using a market basket of post 1L treatments or (as is commonly done including by us) assuming that costs in the progressed state are based on

2L treatment.

# Appendices

# A Population

Figure A1 assess the extent to which the age distribution can be approximated by a normal distribution. The histogram provides an approximation of the empirical age distribution as reported by SEER. The red line is a normally distributed age distribution using the mean and standard deviation from the SEER data. The normal curve tends to match the empirical data, which suggests that the distribution of patient ages in the simulation can be generated from a normal distribution.

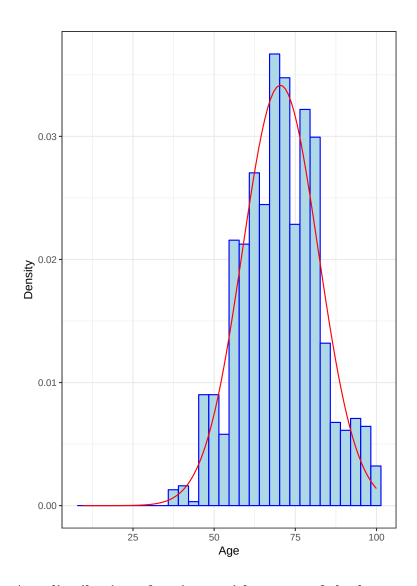


Figure A1: Age distribution of patients with cancer of the lung and bronchus

### B Systematic Literature Review

#### B.1 Treatment effects for transition rates

Details regarding eligibility criteria defining the scope of the studies considered relevant are outlined in Table A1, Table A2, and Table A3.

#### B.2 Utilities

Health state utility values (EQ-5d, HUI2, HUI-3, SF-6D) for the different health states were used in the model as well as disutility estimates associated with adverse events identified by focusing on review or overview studies. Furthermore, published mapping algorithms that allow a non-preference-based measure (generic or disease-specific measure) to be mapped onto a generic preference-based measure are of interest, as well as mapping algorithms between different generic preference-based health state utility values (e.g. between SF-6D and EQ5D) will be searched for. See Appendix Table 4 for study selection criteria.

#### B.3 Resource use, productivity, and cost

Relevant evidence regarding resource use, productivity and cost estimates were identified by means of a review of published CEA modelling studies in NSCLC relevant for the US setting. See Appendix Table 5 for study selection criteria.

#### **B.4** Study identification

Relevant studies were identified by searching the following databases using predefined search strategies: Medical Literature Analysis and Retrieval System Online (MEDLINE), Excerpta Medica database (EMBASE), and Cochrane Central Register of Controlled Trials. The study design filters recommended by the Scottish Intercollegiate Guidelines Network (SIGN) for MEDLINE and EMBASE were used to identify clinical trials. The search included terms related to the generic and brand name of the interventions of interest. The US National Institutes of Health Clinical Trial Registry and EU Clinical Trials Registry were also searched to identify completed clinical trials not yet published to identify any completed or ongoing trials that meet the criteria with results available.

For utility studies, and CEA studies providing evidence on healthcare utilization and costs, the following additional databases were searched in addition: NHS Economic Evaluation Database (NHS EED), Health Economic Evaluations Database (HEED), THe Health Economics Research Centre-maintained mapping algorithm database, and The University of Sheffield's ScHARRHUD database of health utilities' evidence.

Table A1: PICOS criteria for review of treatment effects (metastatic 1L population)

PICOS	Criteria
Population	Adult patients with metastatic non-squamous NSCLC who are EGFR positive and without prior treatment for their disease.
Interventions	The following drugs as monotherapy or in combination with other drugs
	• erlotinib
	• afatinib
	• gefitinib
	• osimertinib
	• dacomitinib
Comparators	
-	• placebo
	• best supportive care (BSC), defined as whichever therapy was judged to be appropriate by the treating physician.
	• any intervention of interest
	• any treatment that facilitates an indirect comparison
Outcomes	
	• OS
	• PFS
	• TTP
Study design	RCTs
Other	English language

Note: Trials where the overall study population is an all-comer population, but subgroup results are reported for the target population of interest are included. A trial or subgroup where the study population is a mixture of non-squamous and squamous patients is included if over 90% is non-squamous.

Where there were gaps in the identified evidence, the search was expanded to include the grey literature, such as reports published by NICE.

#### B.5 Study selection

Two reviewers, working independently, reviewed all abstracts and proceedings identified in each of the searches according to the selection criteria, with the exception of outcome criteria in the efficacy and safety searches, which were only applied during the screening of full-text publications. All studies identified as eligible studies during abstract screening were then screened at a full-text stage by the same two reviewers. The full-text studies identified at this stage were included for data extraction. Following reconciliation between the two investigators, a third reviewer was included to reach consensus for any remaining discrepancies. The process of study identification and selection is summarized with Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) flow diagrams (including reasons for exclusion at both the abstract and full text screening stage).

#### B.6 Data collection

For the clinical studies, two reviewers, working independently, extracted data on study characteristics, interventions, patient characteristics, and outcomes for the final list of included studies. Following reconciliation between the two reviewers, a third reviewer was included to reach consensus on any remaining discrepancies. For all outcomes of interest, information regarding point estimates, variability and uncertainty was obtained. For PFS and OS, hazard ratios and associated information regarding uncertainty were extracted. If results were reported as forest plots, the point estimate and 95\$ percent confidence interval were extracted using DigitizeIt software version 2.1.4 (Bormisoft - Informer Technologies, Inc.). Kaplan Meier curves will also be digitzed using DigitizeIt and the proportion of patients free of the event over time will be extracted and the number of patients at risk over time. Adverse events were collected, if reported. Data collected included number and percent of patients with: any grade 3 or 4 adverse event, any grade 3 or 4 treatment related adverse event, any serious adverse event, any treatment related serious adverse event, and death within 30 days of last treatment. The duration over which safety outcomes were reported were also captured. Discontinuation due to adverse events were also examined.

For utility and cost-effectiveness studies, relevant information for the model was extracted from the source publications and checked by a second-reviewer.

#### **B.7** Limitations

Despite the strengths of the proposed SLR, some limitations are applicable to all SLRs that should be acknowledged. As the evidence base is continually growing, any trials published after the search date will not be captured. Further, any trials that are published close to the search date but are not yet indexed in the databases at the time of the search will not be captured by the search of MEDLINE, EMBASE, and Cochrane Central Register of Controlled Trials. As with any literature

review, this SLR is limited by the use of published data. There is a risk of publication bias as some clinical studies fail to be published while others are published only in abstract form, which presents limited information. Finally, the search and selection were restricted to trials published in English.

Table A2: PICOS criteria for review of treatment effects (metastatic 2L population)

PICOS	Criteria
Population	Adult patients with metastatic non-squamous NSCLC who are EGFR+ positive and who have experienced progression after one line of prior treatment.
	of prior deadlient.
Interventions	The following drugs as monotherapy or in combination with other drugs
	• erlotinib
	• afatinib
	• gefitinib
	• osimertinib
	• dacomitinib
	• nivolumab
	• pembrolizumab
	• atezolizumab
	• bevacizumab
	• platinum-based doublet therapy
Comparators	
	• placebo
	• best supportive care (BSC), defined as whichever therapy was judged to be appropriate by the treating physician.
	• any intervention of interest
	ullet any treatment that facilitates an indirect comparison
Outcomes	
	• OS
	• PFS
	• TTP
Study design	RCTs
Other	English language

Note: Trials where the overall study population is an all-comer population, but subgroup results are reported for the target population of interest are included. A trial or subgroup where the study population is a mixture of non-squamous and squamous patients is included if over 90% is non-squamous.

Table A3: PICOS criteria for review of treatment effects (metastatic 2L+ population)

PICOS	Criteria
Population	Adult patients with metastatic non-squamous NSCLC who are EGFR+ positive and who have experienced progression after two or more prior prior treatments.
Interventions	The following drugs as monotherapy or in combination with other drugs
	• nivolumab
	• pembrolizumab
	• atezolizumab
	• bevacizumab
	• platinum-based doublet the rapy
Comparators	
	• placebo
	• best supportive care (BSC), defined as whichever therapy was judged to be appropriate by the treating physician.
	• any intervention of interest
	$\bullet$ any treatment that facilitates an indirect comparison
Outcomes	
	• OS
	• PFS
	• TTP
Study design	RCTs
Other	English language

Note: Trials where the overall study population is an all-comer population, but subgroup results are reported for the target population of interest are included. A trial or subgroup where the study population is a mixture of non-squamous and squamous patients is included if over 90% is non-squamous.

- C Evidence base, efficacy and safety
- C.1 Kaplan-Meier curves
- C.1.1 1L treatment
- C.1.2 2L treatment
- C.2 Adverse events
- D Network meta-analysis for relative treatment effects with 1L therapy
- D.1 JAGS code
- D.1.1 Fixed effects multistate Weibull and Gompertz network meta-analysis model for estimation of relative treatment effects with each treatment versus gefitinib

```
model{
#timepoints for which PFS and OS data is available
for (i in 1:(Nd-207)){
    # likelihood
    r[i,1:3]~dmulti(p[i,1:3], z[i,1])
    r[i,4:6]~dmulti(p[i,4:6], z[i,2])
    r[i,7:9]~dmulti(p[i,7:9], z[i,3])

p[i,1:3]~ddirch(alpha[])

p[i,4]<-p[i,1]*exp(-(h.sd[i]+h.sp[i])*dt[i,1])
    p[i,5]<-p[i,2]*exp(-h.pd[i]*dt[i,1])+p[i,1]*h.sp[i]*(exp(-(h.sd[i]+h.sp[i])*dt[i,1])-exp(-h.p[i,6]<-1-(p[i,4]+p[i,5])

p[i,7]<-p[i,1]*exp(-(h.sd[i]+h.sp[i])*dt[i,2])
    p[i,8]<-p[i,2]*exp(-h.pd[i]*dt[i,2])+p[i,1]*h.sp[i]*(exp(-(h.sd[i]+h.sp[i])*dt[i,2])-exp(-h.p[i,9]<-1-(p[i,7]+p[i,8])

log(h.sp[i])<- Beta[s[i],a[i],1]+Beta[s[i],a[i],2]*timetrans1[i]</pre>
```

log(h.pd[i])<- Beta[s[i],a[i],4]+Beta[s[i],a[i],5]\*timetrans1[i]

log(h.sd[i])<- Beta[s[i],a[i],3]

```
}
#timepoints for which only OS data is available (from studies with both PFS and OS)
  for (i in (Nd-206):Nd){
  # likelihood
   r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
   p_cond_surv[i] <- exp(-h.OS[i] *dt[i,2])</pre>
   h.OS[i] < -h.sd[i] + h.pd[i]
   log(h.sd[i])<- Beta[s[i],a[i],3]
   log(h.pd[i])<- Beta[s[i],a[i],4]+Beta[s[i],a[i],5]*timetrans1[i]
 }
 #Fixed effects model
for (1 in 1:9){
   for (ll in 1:na[l]){
     Beta[1,11,1] < -mu[1,1] + d[t[1,11],1] - d[t[1,1],1]
     Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
     Beta[1,11,3]<-mu[1,3]
     Beta[1,11,4] < -mu[1,4] + d[t[1,11],3] - d[t[1,1],3]
     Beta[1,11,5]<-mu[1,5]
   }
 }
#FLAURA (Study 10 has a mixed control group, 66% GEF, 34% ERL)
Beta[10,1,1]<-mu[10,1]
Beta[10,1,2]<-mu[10,2]
Beta[10,1,3]<-mu[10,3]
Beta[10,1,4]<-mu[10,4]
Beta[10,1,5]<-mu[10,5]
Beta[10,2,1] < -mu[10,1] + d[5,1] - (0.66*d[1,1] + 0.34*d[2,1])
 Beta[10,2,2] < -mu[10,2] + d[5,2] - (0.66*d[1,2] + 0.34*d[2,2])
Beta[10,2,3]<-mu[10,3]
\mathtt{Beta[10,2,4]} < \mathtt{mu[10,4]} + \mathtt{d[5,3]} - (0.66 * \mathtt{d[1,3]} + 0.34 * \mathtt{d[2,3]})
Beta[10,2,5]<-mu[10,5]
```

```
for (1 in 11:Ns){
    for (ll in 1:na[l]){
      Beta[1,11,1] < -mu[1,1] + d[t[1,11],1] - d[t[1,1],1]
      Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
      Beta[1,11,3]<-mu[1,3]
      Beta[1,11,4] < -mu[1,4] + d[t[1,11],3] - d[t[1,1],3]
      Beta[1,11,5]<-mu[1,5]
    }
  }
  #priors
  for (j in 1:Ns){
    mu[j,1:5] ~ dmnorm(prior_mean_mu[1:5],prior_varcov_mu[,])
  }
  d[1,1]<-0
  d[1,2]<-0
  d[1,3]<-0
  for (k in 2:Nt){
    d[k,1:3] ~ dmnorm(prior_mean_d[1:3],prior_varcov_d[,])
  }
}
```

D.1.2 Fixed effects multistate second order fractional polynomial network meta-analysis model for estimation of relative treatment effects of each treatment versus gefitinib

```
model{
    #timepoints for which PFS and OS data is available
    for (i in 1:(Nd-207)){
        # likelihood
        r[i,1:3]~dmulti(p[i,1:3], z[i,1])
        r[i,4:6]~dmulti(p[i,4:6], z[i,2])
```

```
r[i,7:9]^{dmulti(p[i,7:9], z[i,3])}
            p[i,1:3]~ddirch(alpha[])
            p[i,4]<-p[i,1]*exp(-(h.sd[i]+h.sp[i])*dt[i,1])
             p[i,5] < p[i,2] * exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) - exp(-h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,2] * exp(-h.sd[i]+h.sp[i]) * dt[i,2] * exp(-h.sd[i]+h.sp[i]) * dt[i,2]) + p[i,2] * exp(-h.sd[i]+h.sp[i]) * exp(-h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]) * exp(-h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h
            p[i,6]<-1-(p[i,4]+p[i,5])
            p[i,7]<-p[i,1]*exp(-(h.sd[i]+h.sp[i])*dt[i,2])
             p[i,8] < p[i,2] * exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) - exp(-h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,8] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,8]) + p[i,8] * (exp(-(h.sd[i] + h.sp[i]) * 
            p[i,9]<-1-(p[i,7]+p[i,8])
             log(h.sd[i])<- Beta[s[i],a[i],4]
             log(h.pd[i])<- Beta[s[i],a[i],5]+Beta[s[i],a[i],6]*timetrans1[i]
   }
#timepoints for which only OS data is available (from studies with both PFS and OS)
        for (i in (Nd-206):Nd){
        # likelihood
             r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
            p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
            h.OS[i] < -h.sd[i] + h.pd[i]
             log(h.sd[i])<- Beta[s[i],a[i],4]
             log(h.pd[i])<- Beta[s[i],a[i],5]+Beta[s[i],a[i],6]*timetrans1[i]
    }
    #Fixed effects model
   for (1 in 1:9){
             for (ll in 1:na[l]){
                     \texttt{Beta[1,11,1]} < -\texttt{mu[1,1]} + \texttt{d[t[1,11],1]} - \texttt{d[t[1,1],1]}
                     Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
                     Beta[1,11,3]<-mu[1,3]
                     Beta[1,11,4]<-mu[1,4]
```

```
Beta[1,11,5] < -mu[1,5] + d[t[1,11],3] - d[t[1,1],3]
    Beta[1,11,6]<-mu[1,6]
  }
}
#FLAURA (Study 10 has a mixed control group, 66% GEF, 34% ERL)
Beta[10,1,1]<-mu[10,1]
Beta[10,1,2]<-mu[10,2]
Beta[10,1,3]<-mu[10,3]
Beta[10,1,4]<-mu[10,4]
Beta[10,1,5]<-mu[10,5]
Beta[10,1,6]<-mu[10,6]
Beta[10,2,1] < -mu[10,1] + d[5,1] - (0.66*d[1,1] + 0.34*d[2,1])
Beta[10,2,2] < -mu[10,2] + d[5,2] - (0.66*d[1,2] + 0.34*d[2,2])
Beta[10,2,3]<-mu[10,3]
Beta[10,2,4]<-mu[10,4]
Beta[10,2,5] < -mu[10,5] + d[5,3] - (0.66*d[1,3] + 0.34*d[2,3])
Beta[10,2,6]<-mu[10,6]
for (1 in 11:Ns){
  for (ll in 1:na[l]){
    \texttt{Beta[1,11,1]} < -\texttt{mu[1,1]} + \texttt{d[t[1,11],1]} - \texttt{d[t[1,1],1]}
    Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4]<-mu[1,4]
    \texttt{Beta[1,11,5]} < -\texttt{mu[1,5]} + \texttt{d[t[1,11],3]} - \texttt{d[t[1,1],3]}
    Beta[1,11,6]<-mu[1,6]
  }
}
#priors
for (j in 1:Ns){
  mu[j,1:6] ~ dmnorm(prior_mean_mu[1:6],prior_varcov_mu[,])
}
d[1,1]<-0
```

```
d[1,2]<-0
d[1,3]<-0

for (k in 2:Nt){
   d[k,1:3] ~ dmnorm(prior_mean_d[1:3],prior_varcov_d[,])
}</pre>
```

D.1.3 Random effects multistate Weibull and Gompertz network meta-analysis model for estimation of relative treatment effects of each treatment versus gefitinib

```
model{
            #timepoints for which PFS and OS data is available
            for (i in 1:(Nd-207)){
                         # likelihood
                         r[i,1:3]~dmulti(p[i,1:3], z[i,1])
                         r[i,4:6]~dmulti(p[i,4:6], z[i,2])
                         r[i,7:9]~dmulti(p[i,7:9], z[i,3])
                         p[i,1:3]~ddirch(alpha[])
                          p[i,4] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,1])
                          p[i,5] < -p[i,2] * exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) - exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]
                         p[i,6]<-1-(p[i,4]+p[i,5])
                          p[i,7] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,2])
                          p[i,8] < -p[i,2] * exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) - exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.s
                          p[i,9]<-1-(p[i,7]+p[i,8])
                          log(h.sp[i])<- Beta[s[i],a[i],1]+Beta[s[i],a[i],2]*timetrans1[i]
                          log(h.sd[i])<- Beta[s[i],a[i],3]
                          log(h.pd[i])<- Beta[s[i],a[i],4]+Beta[s[i],a[i],5]*timetrans1[i]
            }
```

#timepoints for which only OS data is available (from studies with both PFS and OS)

```
for (i in (Nd-206):Nd){
 # likelihood
  r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
  p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
  h.OS[i] < -h.sd[i] + h.pd[i]
  log(h.sd[i])<- Beta[s[i],a[i],3]
  log(h.pd[i])<- Beta[s[i],a[i],4]+Beta[s[i],a[i],5]*timetrans1[i]</pre>
}
#Random effects model
for (1 in 1:Ns){
  w[1,1]<-0
  delta[1,1] < -0
}
for (1 in 1:9){
  for (ll in 1:na[l]){
    Beta[1,11,1]<-mu[1,1]+delta[1,11]</pre>
    Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4] < -mu[1,4] + d[t[1,11],3] - d[t[1,1],3]
    Beta[1,11,5]<-mu[1,5]
  }
}
#FLAURA (Study 10 has a mixed control group, 66% GEF, 34% ERL)
Beta[10,1,1]<-mu[10,1]
Beta[10,1,2]<-mu[10,2]
Beta[10,1,3]<-mu[10,3]
Beta[10,1,4]<-mu[10,4]
Beta[10,1,5]<-mu[10,5]
Beta[10,2,1]<-mu[10,1]+delta[10,2]
Beta[10,2,2] < -mu[10,2] + d[5,2] - (0.66*d[1,2] + 0.34*d[2,2])
```

```
Beta[10,2,3]<-mu[10,3]
Beta[10,2,4] < -mu[10,4] + d[5,3] - (0.66*d[1,3] + 0.34*d[2,3])
Beta[10,2,5]<-mu[10,5]
for (1 in 11:Ns){
  for (ll in 1:na[l]){
    Beta[1,11,1]<-mu[1,1]+delta[1,11]</pre>
    Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4] < -mu[1,4] + d[t[1,11],3] - d[t[1,1],3]
    Beta[1,11,5]<-mu[1,5]
  }
}
for (1 in 1:9){
  for (ll in 2:na[l]){
    delta[1,11]~dnorm(md[1,11],taud[1,11])
    md[1,11] < -d[t[1,11],1] -d[t[1,1],1] +sw[1,11]
    w[1,11] \leftarrow (delta[1,11] - d[t[1,11],1] + d[t[1,1],1])
    sw[1,11] <- sum(w[1,1:(11-1)])/(11-1)
    taud[1,11] \leftarrow tau *2*(11-1)/11
  }
}
delta[10,2]~dnorm(md[10,2],taud[10,2])
md[10,2] < -d[5,1] - (0.66*d[1,1]+0.34*d[2,1]) +sw[10,2]
w[10,2] \leftarrow (delta[10,2] - d[5,1] + (0.66*d[1,1]+0.34*d[2,1]))
sw[10,2] <- sum(w[10,1:(2-1)])/(2-1)
taud[10,2] \leftarrow tau *2*(2-1)/2
for (1 in 11:Ns){
  for (ll in 2:na[1]){
    delta[1,11]~dnorm(md[1,11],taud[1,11])
    md[1,11] < -d[t[1,11],1] -d[t[1,1],1] +sw[1,11]
```

```
w[1,11] \leftarrow (delta[1,11] - d[t[1,11],1] + d[t[1,1],1])
      sw[1,11] <- sum(w[1,1:(11-1)])/(11-1)
      taud[1,11] <- tau *2*(11-1)/11
    }
  }
  #priors
  for (j in 1:Ns){
    mu[j,1:5] ~ dmnorm(prior_mean_mu[1:5],prior_varcov_mu[,])
  }
  d[1,1]<-0
  d[1,2]<-0
  d[1,3]<-0
  for (k in 2:Nt){
    d[k,1:3] ~ dmnorm(prior_mean_d[1:3],prior_varcov_d[,])
  }
  sd~dunif(0,2)
  tau < -1/(sd*sd)
}
```

D.1.4 Random effects multistate second order fractional polynomial network metaanalysis model for estimation of relative treatment effects of each treatment versus gefitinib

```
model{
    #timepoints for which PFS and OS data is available
    for (i in 1:(Nd-207)){
        # likelihood
        r[i,1:3]~dmulti(p[i,1:3], z[i,1])
        r[i,4:6]~dmulti(p[i,4:6], z[i,2])
        r[i,7:9]~dmulti(p[i,7:9], z[i,3])
```

```
p[i,1:3]~ddirch(alpha[])
              p[i,4] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,1])
              p[i,5] < -p[i,2] * exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) - exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]
              p[i,6]<-1-(p[i,4]+p[i,5])
              p[i,7] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,2])
              p[i,8] < -p[i,2] * exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) - exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.s
              p[i,9]<-1-(p[i,7]+p[i,8])
              log(h.sp[i])<- Beta[s[i],a[i],1]+Beta[s[i],a[i],2]*timetrans1[i]+Beta[s[i],a[i],3]*timetrans
              log(h.sd[i])<- Beta[s[i],a[i],4]
              log(h.pd[i])<- Beta[s[i],a[i],5]+Beta[s[i],a[i],6]*timetrans1[i]
    }
#timepoints for which only OS data is available (from studies with both PFS and OS)
         for (i in (Nd-206):Nd){
         # likelihood
              r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
              p_cond_surv[i] <- exp(-h.OS[i] *dt[i,2])</pre>
              h.OS[i] < -h.sd[i] + h.pd[i]
              log(h.sd[i])<- Beta[s[i],a[i],4]
              log(h.pd[i])<- Beta[s[i],a[i],5]+Beta[s[i],a[i],6]*timetrans1[i]
    }
    #Random effects model
    for (1 in 1:Ns){
              w[1,1]<-0
              delta[1,1] < -0
    }
    for (1 in 1:9){
              for (ll in 1:na[l]){
```

```
Beta[1,11,1]<-mu[1,1]+delta[1,11]
    Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4]<-mu[1,4]
    Beta[1,11,5] < -mu[1,5] + d[t[1,11],3] - d[t[1,1],3]
    Beta[1,11,6]<-mu[1,6]
  }
}
#FLAURA (Study 10 has a mixed control group, 66% GEF, 34% ERL)
Beta[10,1,1]<-mu[10,1]
Beta[10,1,2]<-mu[10,2]
Beta[10,1,3]<-mu[10,3]
Beta[10,1,4]<-mu[10,4]
Beta[10,1,5]<-mu[10,5]
Beta[10,1,6]<-mu[10,6]
Beta[10,2,1]<-mu[10,1]+delta[10,2]
Beta[10,2,2] < -mu[10,2] + d[5,2] - (0.66*d[1,2] + 0.34*d[2,2])
Beta[10,2,3]<-mu[10,3]
Beta[10,2,4]<-mu[10,4]
Beta[10,2,5] < -mu[10,5] + d[5,3] - (0.66*d[1,3]+0.34*d[2,3])
Beta[10,2,6]<-mu[10,6]
for (1 in 11:Ns){
  for (ll in 1:na[l]){
    Beta[1,11,1]<-mu[1,1]+delta[1,11]
    Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4]<-mu[1,4]
    \texttt{Beta[1,11,5]} < -\texttt{mu[1,5]} + \texttt{d[t[1,11],3]} - \texttt{d[t[1,1],3]}
    Beta[1,11,6]<-mu[1,6]
  }
}
```

```
for (1 in 1:9){
  for (ll in 2:na[l]){
    delta[1,11]~dnorm(md[1,11],taud[1,11])
    md[1,11] < -d[t[1,11],1] -d[t[1,1],1] +sw[1,11]
    w[1,11] \leftarrow (delta[1,11] - d[t[1,11],1] + d[t[1,1],1])
    sw[1,11] <- sum(w[1,1:(11-1)])/(11-1)
    taud[1,11] <- tau *2*(11-1)/11
  }
}
delta[10,2]~dnorm(md[10,2],taud[10,2])
md[10,2] < -d[5,1] - (0.66*d[1,1]+0.34*d[2,1]) +sw[10,2]
w[10,2] \leftarrow (delta[10,2] - d[5,1] + (0.66*d[1,1]+0.34*d[2,1]))
sw[10,2] <- sum(w[10,1:(2-1)])/(2-1)
taud[10,2] \leftarrow tau *2*(2-1)/2
for (1 in 11:Ns){
  for (ll in 2:na[1]){
    delta[1,11]~dnorm(md[1,11],taud[1,11])
    md[1,11] < -d[t[1,11],1] -d[t[1,1],1] +sw[1,11]
    w[1,11] \leftarrow (delta[1,11] - d[t[1,11],1] + d[t[1,1],1])
    sw[1,11] <- sum(w[1,1:(11-1)])/(11-1)
    taud[1,11] <- tau *2*(11-1)/11
 }
}
#priors
for (j in 1:Ns){
 mu[j,1:6] ~ dmnorm(prior_mean_mu[1:6],prior_varcov_mu[,])
}
d[1,1]<-0
d[1,2]<-0
d[1,3]<-0
```

```
for (k in 2:Nt){
   d[k,1:3] ~ dmnorm(prior_mean_d[1:3],prior_varcov_d[,])
}

sd~dunif(0,2)
tau<-1/(sd*sd)
}</pre>
```

D.1.5 Fixed effects multistate Weibull and Gompertz network meta-analysis model for estimation of relative treatment effects of each treatment versus gefitinib; no treatment effect on PD transitions

```
model{
              #timepoints for which PFS and OS data is available
             for (i in 1:(Nd-207)){
                          # likelihood
                          r[i,1:3]~dmulti(p[i,1:3], z[i,1])
                          r[i,4:6]~dmulti(p[i,4:6], z[i,2])
                          r[i,7:9]~dmulti(p[i,7:9], z[i,3])
                          p[i,1:3]~ddirch(alpha[])
                          p[i,4] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,1])
                           p[i,5] < -p[i,2] * exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) - exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]
                           p[i,6]<-1-(p[i,4]+p[i,5])
                          p[i,7] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,2])
                           p[i,8] < -p[i,2] * exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) - exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.s
                           p[i,9]<-1-(p[i,7]+p[i,8])
                           log(h.sp[i])<- Beta[s[i],a[i],1]+Beta[s[i],a[i],2]*timetrans1[i]
                           log(h.sd[i])<- Beta[s[i],a[i],3]
                           log(h.pd[i])<- Beta[s[i],a[i],4]+Beta[s[i],a[i],5]*timetrans1[i]
```

}

```
#timepoints for which only OS data is available (from studies with both PFS and OS)
  for (i in (Nd-206):Nd){
  # likelihood
   r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
   p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
   h.OS[i] < -h.sd[i] + h.pd[i]
   log(h.sd[i])<- Beta[s[i],a[i],3]
   log(h.pd[i])<- Beta[s[i],a[i],4]+Beta[s[i],a[i],5]*timetrans1[i]
}
#Fixed effects model
for (1 in 1:9){
   for (ll in 1:na[l]){
     Beta[1,11,1] < -mu[1,1] + d[t[1,11],1] - d[t[1,1],1]
     Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
     Beta[1,11,3]<-mu[1,3]
     Beta[1,11,4]<-mu[1,4]
     Beta[1,11,5]<-mu[1,5]
   }
}
#FLAURA (Study 10 has a mixed control group, 66% GEF, 34% ERL)
Beta[10,1,1]<-mu[10,1]
Beta[10,1,2]<-mu[10,2]
Beta[10,1,3]<-mu[10,3]
Beta[10,1,4]<-mu[10,4]
Beta[10,1,5]<-mu[10,5]
Beta[10,2,1] < -mu[10,1] + d[5,1] - (0.66*d[1,1] + 0.34*d[2,1])
Beta[10,2,2] < -mu[10,2] + d[5,2] - (0.66*d[1,2] + 0.34*d[2,2])
Beta[10,2,3]<-mu[10,3]
Beta[10,2,4]<-mu[10,4]
Beta[10,2,5]<-mu[10,5]
```

```
for (1 in 11:Ns){
  for (ll in 1:na[l]){
    Beta[1,11,1] < -mu[1,1] + d[t[1,11],1] - d[t[1,1],1]
    Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4]<-mu[1,4]
    Beta[1,11,5]<-mu[1,5]
 }
}
#priors
for (j in 1:Ns){
  mu[j,1:5] ~ dmnorm(prior_mean_mu[1:5],prior_varcov_mu[,])
}
d[1,1]<-0
d[1,2]<-0
for (k in 2:Nt){
  d[k,1:2] ~ dmnorm(prior_mean_d[1:2],prior_varcov_d[,])
}
```

D.1.6 Fixed effects multistate second order fractional polynomial network meta-analysis model for estimation of relative treatment effects of each treatment versus gefitinib; no treatment effect on PD transitions

```
model{
    #timepoints for which PFS and OS data is available
    for (i in 1:(Nd-207)){
        # likelihood
        r[i,1:3]~dmulti(p[i,1:3], z[i,1])
        r[i,4:6]~dmulti(p[i,4:6], z[i,2])
        r[i,7:9]~dmulti(p[i,7:9], z[i,3])

        p[i,1:3]~ddirch(alpha[])
```

```
p[i,4] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,1])
             p[i,5] < -p[i,2] * exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) - exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]
            p[i,6]<-1-(p[i,4]+p[i,5])
            p[i,7] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,2])
            p[i,8] < p[i,2] * exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) - exp(-h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,2] * exp(-h.sd[i] + h.sp[i]) * exp(-h.sd[i] + h.sp[i] + h.sp[i]) * exp(-h.sd[i] + h.sp[i] + h.sp[i] * exp(-h.sd[i] + h.sp[i] + h.sp[
             p[i,9]<-1-(p[i,7]+p[i,8])
             log(h.sd[i])<- Beta[s[i],a[i],4]
             log(h.pd[i])<- Beta[s[i],a[i],5]+Beta[s[i],a[i],6]*timetrans1[i]
   }
#timepoints for which only OS data is available (from studies with both PFS and OS)
        for (i in (Nd-206):Nd){
        # likelihood
            r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
            p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
             h.OS[i] < -h.sd[i] + h.pd[i]
             log(h.sd[i])<- Beta[s[i],a[i],4]
             log(h.pd[i])<- Beta[s[i],a[i],5]+Beta[s[i],a[i],6]*timetrans1[i]
    }
   #Fixed effects model
   for (1 in 1:9){
            for (ll in 1:na[l]){
                     Beta[1,11,1] < -mu[1,1] + d[t[1,11],1] - d[t[1,1],1]
                     Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
                     Beta[1,11,3]<-mu[1,3]
                     Beta[1,11,4]<-mu[1,4]
                     Beta[1,11,5]<-mu[1,5]
                     Beta[1,11,6]<-mu[1,6]
             }
```

```
}
#FLAURA (Study 10 has a mixed control group, 66% GEF, 34% ERL)
Beta[10,1,1]<-mu[10,1]
Beta[10,1,2]<-mu[10,2]
Beta[10,1,3]<-mu[10,3]
Beta[10,1,4]<-mu[10,4]
Beta[10,1,5]<-mu[10,5]
Beta[10,1,6]<-mu[10,6]
Beta[10,2,1] < -mu[10,1] + d[5,1] - (0.66*d[1,1] + 0.34*d[2,1])
Beta[10,2,2] < -mu[10,2] + d[5,2] - (0.66*d[1,2] + 0.34*d[2,2])
Beta[10,2,3]<-mu[10,3]
Beta[10,2,4]<-mu[10,4]
Beta[10,2,5]<-mu[10,5]
Beta[10,2,6]<-mu[10,6]
for (1 in 11:Ns){
  for (ll in 1:na[l]){
    Beta[1,11,1] < -mu[1,1] + d[t[1,11],1] - d[t[1,1],1]
    Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4]<-mu[1,4]
    Beta[1,11,5]<-mu[1,5]
    Beta[1,11,6]<-mu[1,6]
  }
}
#priors
for (j in 1:Ns){
  mu[j,1:6] ~ dmnorm(prior_mean_mu[1:6],prior_varcov_mu[,])
}
d[1,1]<-0
d[1,2]<-0
```

```
for (k in 2:Nt){
   d[k,1:2] ~ dmnorm(prior_mean_d[1:2],prior_varcov_d[,])
}
```

D.1.7 Random effects multistate Weibull and Gompertz network meta-analysis model for estimation of relative treatment effects of each treatment versus gefitinib; no treatment effect on PD transitions

```
model{
           #timepoints for which PFS and OS data is available
           for (i in 1:(Nd-207)){
                       # likelihood
                       r[i,1:3]~dmulti(p[i,1:3], z[i,1])
                       r[i,4:6]~dmulti(p[i,4:6], z[i,2])
                       r[i,7:9]~dmulti(p[i,7:9], z[i,3])
                       p[i,1:3]~ddirch(alpha[])
                       p[i,4] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,1])
                       p[i,5] < -p[i,2] * exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) - exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]) * (exp(-(h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]
                       p[i,6]<-1-(p[i,4]+p[i,5])
                       p[i,7] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,2])
                       p[i,8] < -p[i,2] * exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) - exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.s
                       p[i,9]<-1-(p[i,7]+p[i,8])
                       log(h.sp[i])<- Beta[s[i],a[i],1]+Beta[s[i],a[i],2]*timetrans1[i]
                       log(h.sd[i])<- Beta[s[i],a[i],3]
                       log(h.pd[i])<- Beta[s[i],a[i],4]+Beta[s[i],a[i],5]*timetrans1[i]
             }
      #timepoints for which only OS data is available (from studies with both PFS and OS)
                 for (i in (Nd-206):Nd){
                 # likelihood
```

```
r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
  p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
  h.OS[i] < -h.sd[i] + h.pd[i]
  log(h.sd[i])<- Beta[s[i],a[i],3]
  log(h.pd[i])<- Beta[s[i],a[i],4]+Beta[s[i],a[i],5]*timetrans1[i]
}
#Random effects model
for (1 in 1:Ns){
  w[1,1]<-0
  delta[1,1] < -0
}
for (1 in 1:9){
  for (ll in 1:na[l]){
    Beta[1,11,1]<-mu[1,1]+delta[1,11]
    Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4]<-mu[1,4]
    Beta[1,11,5]<-mu[1,5]
  }
}
#FLAURA (Study 10 has a mixed control group, 66% GEF, 34% ERL)
Beta[10,1,1]<-mu[10,1]
Beta[10,1,2]<-mu[10,2]
Beta[10,1,3]<-mu[10,3]
Beta[10,1,4]<-mu[10,4]
Beta[10,1,5]<-mu[10,5]
Beta[10,2,1]<-mu[10,1]+delta[10,2]
Beta[10,2,2] < -mu[10,2] + d[5,2] - (0.66*d[1,2] + 0.34*d[2,2])
Beta[10,2,3]<-mu[10,3]
Beta[10,2,4]<-mu[10,4]
```

```
Beta[10,2,5]<-mu[10,5]
for (1 in 11:Ns){
  for (ll in 1:na[l]){
    Beta[1,11,1]<-mu[1,1]+delta[1,11]
    Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4]<-mu[1,4]
    Beta[1,11,5]<-mu[1,5]
  }
}
for (1 in 1:9){
  for (ll in 2:na[l]){
    delta[1,11]~dnorm(md[1,11],taud[1,11])
    md[1,11] < -d[t[1,11],1] -d[t[1,1],1] +sw[1,11]
    w[1,11] \leftarrow (delta[1,11] - d[t[1,11],1] + d[t[1,1],1])
    sw[1,11] <- sum(w[1,1:(11-1)])/(11-1)
    taud[1,11] <- tau *2*(11-1)/11
  }
}
delta[10,2]~dnorm(md[10,2],taud[10,2])
md[10,2] < -d[5,1] - (0.66*d[1,1]+0.34*d[2,1]) +sw[10,2]
w[10,2] \leftarrow (delta[10,2] - d[5,1] + (0.66*d[1,1]+0.34*d[2,1]))
sw[10,2] <- sum(w[10,1:(2-1)])/(2-1)
taud[10,2] \leftarrow tau *2*(2-1)/2
for (1 in 11:Ns){
  for (ll in 2:na[l]){
    delta[1,11]~dnorm(md[1,11],taud[1,11])
    md[1,11] < -d[t[1,11],1] -d[t[1,1],1] +sw[1,11]
    w[1,11] \leftarrow (delta[1,11] - d[t[1,11],1] + d[t[1,1],1])
```

```
sw[1,11] <- sum(w[1,1:(11-1)])/(11-1)
      taud[1,11] <- tau *2*(11-1)/11
    }
  }
  #priors
  for (j in 1:Ns){
   mu[j,1:5] ~ dmnorm(prior_mean_mu[1:5],prior_varcov_mu[,])
  }
  d[1,1]<-0
  d[1,2]<-0
  for (k in 2:Nt){
    d[k,1:2] ~ dmnorm(prior_mean_d[1:2],prior_varcov_d[,])
  }
  sd~dunif(0,2)
  tau < -1/(sd*sd)
}
```

D.1.8 Random effects multistate second order fractional polynomial network metaanalysis model for estimation of relative treatment effects of each treatment versus gesitinib; no treatment effect on PD transitions

```
model{
    #timepoints for which PFS and OS data is available
    for (i in 1:(Nd-207)){
        # likelihood
        r[i,1:3]~dmulti(p[i,1:3], z[i,1])
        r[i,4:6]~dmulti(p[i,4:6], z[i,2])
        r[i,7:9]~dmulti(p[i,7:9], z[i,3])
```

```
p[i,1:3]~ddirch(alpha[])
             p[i,4]<-p[i,1]*exp(-(h.sd[i]+h.sp[i])*dt[i,1])
              p[i,5] < p[i,2] * exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i]+h.sp[i]) * dt[i,1]) - exp(-h.sd[i]+h.sp[i]) * dt[i,1]) + p[i,2] * exp(-h.sd[i]+h.sp[i]) * dt[i,2] * exp(-h.sd[i]+h.sp[i]) * dt[i,2]) + p[i,2] * exp(-h.sd[i]+h.sp[i]) * exp(-h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]) * exp(-h.sd[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h.sp[i]+h
              p[i,6]<-1-(p[i,4]+p[i,5])
              p[i,7] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,2])
              p[i,8] < -p[i,2] * exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) - exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h
              p[i,9]<-1-(p[i,7]+p[i,8])
              log(h.sd[i])<- Beta[s[i],a[i],4]
              log(h.pd[i])<- Beta[s[i],a[i],5]+Beta[s[i],a[i],6]*timetrans1[i]
   }
#timepoints for which only OS data is available (from studies with both PFS and OS)
        for (i in (Nd-206):Nd){
        # likelihood
             r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
             p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
             h.OS[i] < -h.sd[i] + h.pd[i]
              log(h.sd[i])<- Beta[s[i],a[i],4]
              log(h.pd[i])<- Beta[s[i],a[i],5]+Beta[s[i],a[i],6]*timetrans1[i]
    }
   #Random effects model
   for (1 in 1:Ns){
             w[1,1]<-0
             delta[1,1] < -0
   }
   for (l in 1:9){
              for (ll in 1:na[l]){
                       Beta[1,11,1]<-mu[1,1]+delta[1,11]
```

```
Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4]<-mu[1,4]
    Beta[1,11,5]<-mu[1,5]
    Beta[1,11,6]<-mu[1,6]
  }
}
#FLAURA (Study 10 has a mixed control group, 66% GEF, 34% ERL)
Beta[10,1,1]<-mu[10,1]
Beta[10,1,2]<-mu[10,2]
Beta[10,1,3]<-mu[10,3]
Beta[10,1,4]<-mu[10,4]
Beta[10,1,5]<-mu[10,5]
Beta[10,1,6]<-mu[10,6]
Beta[10,2,1]<-mu[10,1]+delta[10,2]
Beta[10,2,2] < -mu[10,2] + d[5,2] - (0.66*d[1,2] + 0.34*d[2,2])
Beta[10,2,3]<-mu[10,3]
Beta[10,2,4]<-mu[10,4]
Beta[10,2,5]<-mu[10,5]
Beta[10,2,6]<-mu[10,6]
for (1 in 11:Ns){
  for (ll in 1:na[l]){
    Beta[1,11,1]<-mu[1,1]+delta[1,11]
    Beta[1,11,2] < -mu[1,2] + d[t[1,11],2] - d[t[1,1],2]
    Beta[1,11,3]<-mu[1,3]
    Beta[1,11,4]<-mu[1,4]
    Beta[1,11,5]<-mu[1,5]
    Beta[1,11,6]<-mu[1,6]
  }
}
```

```
for (1 in 1:9){
  for (ll in 2:na[l]){
    delta[1,11]~dnorm(md[1,11],taud[1,11])
    md[1,11] < -d[t[1,11],1] - d[t[1,1],1] +sw[1,11]
    w[1,11] \leftarrow (delta[1,11] - d[t[1,11],1] + d[t[1,1],1])
    sw[1,11] <- sum(w[1,1:(11-1)])/(11-1)
    taud[1,11] <- tau *2*(11-1)/11
  }
}
delta[10,2]~dnorm(md[10,2],taud[10,2])
md[10,2] < -d[5,1] - (0.66*d[1,1]+0.34*d[2,1]) +sw[10,2]
w[10,2] \leftarrow (delta[10,2] - d[5,1] + (0.66*d[1,1]+0.34*d[2,1]))
sw[10,2] <- sum(w[10,1:(2-1)])/(2-1)
taud[10,2] \leftarrow tau *2*(2-1)/2
for (1 in 11:Ns){
  for (ll in 2:na[l]){
    delta[1,11]~dnorm(md[1,11],taud[1,11])
    md[1,11] < -d[t[1,11],1] - d[t[1,1],1] +sw[1,11]
    w[1,11] \leftarrow (delta[1,11] - d[t[1,11],1] + d[t[1,1],1])
    sw[1,11] <- sum(w[1,1:(11-1)])/(11-1)
    taud[1,11] <- tau *2*(11-1)/11
  }
}
#priors
for (j in 1:Ns){
 mu[j,1:6] ~ dmnorm(prior_mean_mu[1:6],prior_varcov_mu[,])
}
d[1,1]<-0
d[1,2]<-0
for (k in 2:Nt){
```

```
d[k,1:2] ~ dmnorm(prior_mean_d[1:2],prior_varcov_d[,])
}

sd~dunif(0,2)
tau<-1/(sd*sd)
}</pre>
```

- D.2 Model parameter estimates
- E Meta-analysis for absolute effects with 1L reference treatment
- E.1 JAGS code

log(h.sd[i])<- MU[3]

E.1.1 Fixed effects multistate Weibull and Gompertz meta-analysis model for estimation of absolute effects with 1L gefitinib

```
model{
#timepoints for which PFS and OS data is available
for (i in 1:(Nd-38)){
    # likelihood
    r[i,1:3]~dmulti(p[i,1:3], z[i,1])
    r[i,4:6]~dmulti(p[i,4:6], z[i,2])
    r[i,7:9]~dmulti(p[i,7:9], z[i,3])

p[i,1:3]~ddirch(alpha[])

p[i,4]<-p[i,1]*exp(-(h.sd[i]+h.sp[i])*dt[i,1])
    p[i,5]<-p[i,2]*exp(-h.pd[i]*dt[i,1])+p[i,1]*h.sp[i]*(exp(-(h.sd[i]+h.sp[i])*dt[i,1])-exp(-h.p[i,6]<-1-(p[i,4]+p[i,5])

p[i,7]<-p[i,1]*exp(-(h.sd[i]+h.sp[i])*dt[i,2])
    p[i,8]<-p[i,2]*exp(-h.pd[i]*dt[i,2])+p[i,1]*h.sp[i]*(exp(-(h.sd[i]+h.sp[i])*dt[i,2])-exp(-h.p[i,9]<-1-(p[i,7]+p[i,8])

log(h.sp[i])<- MU[1]+MU[2]*timetrans1[i]</pre>
```

```
log(h.pd[i])<- MU[4]+MU[5]*timetrans1[i]
}

#timepoints for which only OS data is available (from studies with both PFS and OS)
for (i in (Nd-37):Nd){
    # likelihood
    r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])

    p_cond_surv[i]<-exp(-h.OS[i]*dt[i,2])
    h.OS[i]<-h.sd[i]+h.pd[i]

log(h.sd[i])<- MU[3]
log(h.pd[i])<- MU[4]+MU[5]*timetrans1[i]
}

#priors
    MU[1:5] ~ dmnorm(prior_mean_mu[1:5],prior_varcov_mu[,])</pre>
```

# E.1.2 Fixed effects multistate 2nd order fractional polynomial meta-analysis model for estimation of absolute effects with 1L gefitinib

```
model{
    #timepoints for which PFS and OS data is available
    for (i in 1:(Nd-38)){
        # likelihood
        r[i,1:3]~dmulti(p[i,1:3], z[i,1])
        r[i,4:6]~dmulti(p[i,4:6], z[i,2])
        r[i,7:9]~dmulti(p[i,7:9], z[i,3])

        p[i,1:3]~ddirch(alpha[])

        p[i,4]<-p[i,1]*exp(-(h.sd[i]+h.sp[i])*dt[i,1])
        p[i,5]<-p[i,2]*exp(-h.pd[i]*dt[i,1])+p[i,1]*h.sp[i]*(exp(-(h.sd[i]+h.sp[i])*dt[i,1])-exp(-h.p[i,6]<-1-(p[i,4]+p[i,5])</pre>
```

```
p[i,7] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,2])
           p[i,8] < -p[i,2] * exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) - exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.s
           p[i,9]<-1-(p[i,7]+p[i,8])
           log(h.sp[i])<- MU[1]+MU[2]*timetrans1[i]+MU[3]*timetrans2[i]</pre>
           log(h.sd[i])<- MU[4]
           log(h.pd[i])<- MU[5]+MU[6]*timetrans1[i]</pre>
   }
#timepoints for which only OS data is available (from studies with both PFS and OS)
       for (i in (Nd-37):Nd){
       # likelihood
           r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
           p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
           h.OS[i] < -h.sd[i] + h.pd[i]
           log(h.sd[i])<- MU[4]
           log(h.pd[i])<- MU[5]+MU[6]*timetrans1[i]</pre>
   }
   #priors
           MU[1:6] ~ dmnorm(prior_mean_mu[1:6],prior_varcov_mu[,])
```

}

#### E.2 Model parameter estimates

## F Analysis of absolute effects with 2L therapy

#### F.1 JAGS code

F.1.1 Fixed effects multistate Weibull and Gompertz model for estimation of absolute effects with 2L osimertinib

```
model{
 #timepoints for which only OS data is available (from studies with both PFS and OS)
   for (i in 1:(Nd-6)){
   # likelihood
    r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
    p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
    h.OS[i] < -h.sd[i] + h.pd[i]
    log(h.sd[i])<- MU[3]
    log(h.pd[i])<- MU[4]+MU[5]*timetrans1[i]</pre>
 }
  #timepoints for which only PFS data is available
 for (i in (Nd-5):Nd){
    # likelihood
    r_cond_pfs[i]~dbinom(p_cond_pfs[i], z_cond_pfs[i])
    p_cond_pfs[i] <-exp(-h.PFS[i]*dt[i,2])</pre>
    h.PFS[i]<-h.sp[i]+h.sd[i]
    log(h.sp[i])<- MU[1]+MU[2]*timetrans1[i]</pre>
    log(h.sd[i])<- MU[3]
      }
  #priors
    MU[1:5] ~ dmnorm(prior_mean_mu[1:5],prior_varcov_mu[,])
```

F.1.2 Fixed effects multistate 2nd order fractional polynomial model for estimation of absolute effects with 2L osimertinib

```
model{
 #timepoints for which only OS data is available (from studies with both PFS and OS)
   for (i in 1:(Nd-6)){
   # likelihood
    r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
    p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
    h.OS[i] < -h.sd[i] + h.pd[i]
    log(h.sd[i])<- MU[4]
    log(h.pd[i])<- MU[5]+MU[6]*timetrans1[i]</pre>
  }
  #timepoints for which only PFS data is available
  for (i in (Nd-5):Nd){
    # likelihood
    r_cond_pfs[i]~dbinom(p_cond_pfs[i], z_cond_pfs[i])
    p_cond_pfs[i] <-exp(-h.PFS[i]*dt[i,2])</pre>
    h.PFS[i]<-h.sp[i]+h.sd[i]
    log(h.sp[i])<- MU[1]+MU[2]*timetrans1[i]+MU[3]*timetrans2[i]</pre>
    log(h.sd[i])<- MU[4]
      }
  #priors
  MU[1:6] ~ dmnorm(prior_mean_mu[1:6],prior_varcov_mu[,])
}
```

# F.1.3 Fixed effects multistate Weibull and Gompertz model for estimation of absolute effects with 2L PBDC

```
model{
          #timepoints for which PFS and OS data is available
         for (i in 1:(Nd-14)){
                    # likelihood
                    r[i,1:3]~dmulti(p[i,1:3], z[i,1])
                    r[i,4:6]~dmulti(p[i,4:6], z[i,2])
                    r[i,7:9]~dmulti(p[i,7:9], z[i,3])
                    p[i,1:3]~ddirch(alpha[])
                    p[i,4] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,1])
                    p[i,5] < -p[i,2] * exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) - exp(-h.pd[i] * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.sp[i]) * dt[i,1]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-
                    p[i,6]<-1-(p[i,4]+p[i,5])
                    p[i,7]<-p[i,1]*exp(-(h.sd[i]+h.sp[i])*dt[i,2])
                    p[i,8] < -p[i,2] * exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) - exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.s
                    p[i,9]<-1-(p[i,7]+p[i,8])
                    log(h.sp[i])<- MU[1]+MU[2]*timetrans1[i]</pre>
                    log(h.sd[i])<- MU[3]
                    log(h.pd[i])<- MU[4]+MU[5]*timetrans1[i]</pre>
         }
     #timepoints for which only OS data is available (from studies with both PFS and OS)
               for (i in (Nd-13):(Nd-6)){
               # likelihood
                    r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
                    p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
                    h.OS[i] < -h.sd[i] + h.pd[i]
                    log(h.sd[i])<- MU[3]
                    log(h.pd[i])<- MU[4]+MU[5]*timetrans1[i]</pre>
          }
```

```
#timepoints for which only PFS data is available
for (i in (Nd-5):Nd){
    # likelihood
    r_cond_pfs[i]~dbinom(p_cond_pfs[i], z_cond_pfs[i])

    p_cond_pfs[i]<-exp(-h.PFS[i]*dt[i,2])
    h.PFS[i]<-h.sp[i]+h.sd[i]

log(h.sp[i])<- MU[1]+MU[2]*timetrans1[i]
log(h.sd[i])<- MU[3]
    }

#priors
    MU[1:5] ~ dmnorm(prior_mean_mu[1:5],prior_varcov_mu[,])
}</pre>
```

# F.1.4 Fixed effects multistate 2nd order fractional polynomial model for estimation of absolute effects with 2L PBDC

```
model{
    #timepoints for which PFS and OS data is available
    for (i in 1:(Nd-14)){
        # likelihood
        r[i,1:3]~dmulti(p[i,1:3], z[i,1])
        r[i,4:6]~dmulti(p[i,4:6], z[i,2])
        r[i,7:9]~dmulti(p[i,7:9], z[i,3])

        p[i,1:3]~ddirch(alpha[])

        p[i,4]<-p[i,1]*exp(-(h.sd[i]+h.sp[i])*dt[i,1])
        p[i,5]<-p[i,2]*exp(-h.pd[i]*dt[i,1])+p[i,1]*h.sp[i]*(exp(-(h.sd[i]+h.sp[i])*dt[i,1])-exp(-h.pl[i,6]<-1-(p[i,4]+p[i,5])</pre>
```

```
p[i,7] < -p[i,1] * exp(-(h.sd[i]+h.sp[i])*dt[i,2])
         p[i,8] < -p[i,2] * exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) - exp(-h.pd[i] * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * h.sp[i] * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,1] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.sp[i]) * dt[i,2]) + p[i,2] * (exp(-(h.sd[i] + h.sp[i]) * (exp(-(h.sd[i] + h.s
         p[i,9]<-1-(p[i,7]+p[i,8])
         log(h.sp[i])<- MU[1]+MU[2]*timetrans1[i]+MU[3]*timetrans2[i]</pre>
         log(h.sd[i])<- MU[4]
         log(h.pd[i])<- MU[5]+MU[6]*timetrans1[i]</pre>
  }
#timepoints for which only OS data is available (from studies with both PFS and OS)
     for (i in (Nd-13):(Nd-6)){
     # likelihood
        r_cond_surv[i]~dbinom(p_cond_surv[i], z_cond_surv[i])
        p_cond_surv[i] <-exp(-h.OS[i]*dt[i,2])</pre>
        h.OS[i] < -h.sd[i] + h.pd[i]
         log(h.sd[i])<- MU[4]
         log(h.pd[i])<- MU[5]+MU[6]*timetrans1[i]</pre>
  }
  #timepoints for which only PFS data is available
  for (i in (Nd-5):Nd){
        # likelihood
        r_cond_pfs[i]~dbinom(p_cond_pfs[i], z_cond_pfs[i])
         p_cond_pfs[i] <-exp(-h.PFS[i]*dt[i,2])</pre>
         h.PFS[i]<-h.sp[i]+h.sd[i]
         log(h.sp[i])<- MU[1]+MU[2]*timetrans1[i]+MU[3]*timetrans2[i]</pre>
         log(h.sd[i])<- MU[4]
               }
  #priors
  MU[1:6] ~ dmnorm(prior_mean_mu[1:6],prior_varcov_mu[,])
```

### F.2 Model parameter estimates

## G Network meta-analysis of adverse events

### G.1 JAGS code

#### G.1.1 Alt increase

# Priors d[1]<-0

```
model{
  # Binomial likelihood, logit link
  # Model for relative treatment effects
    for(i in 1:(ns-1)){
                                         # LOOP THROUGH STUDIES
      for (k in 1:na[i]) {
                                        # LOOP THROUGH ARMS
        r[i,k] ~ dbin(p[i,k],n[i,k]) # binomial likelihood
        logit(p[i,k]) \leftarrow mu[i] + d[t[i,k]] - d[t[i,1]] # model for linear predictor
      }
    }
                                          # LOOP THROUGH ARMS OF LAST STUDY (FLAURA)
      for (k in 1:na[ns]) {
        r[ns,k] ~ dbin(p[ns,k],n[ns,k]) # binomial likelihood
        logit(p[ns,k]) \leftarrow mu[ns] + d[t[ns,k]] - (0.66*d[1]+0.34*d[2]) # model for linear predictions.
      }
  # Exchangeable relative treatment effects for TKIs
    for (k in 2:5){ d[k] ~ dnorm(d.TKI,d.TKI.prec) }
  # Random effects model for absolute effects with GEF
   mu[1] ~ dnorm(MU,MU.prec)
   mu[3] ~ dnorm(MU,MU.prec)
   mu[4] ~ dnorm(MU,MU.prec)
   mu[8] ~ dnorm(MU,MU.prec)
   mu[9] ~ dnorm(MU,MU.prec)
```

# treatment effect is zero for reference treatm

```
for (k \text{ in } 6:\text{nt})\{ d[k] \sim dnorm(0,.001) \} # vague priors for treatment effects
  d.TKI ~ dnorm(0,.001)
  d.TKI.sd ~ dunif(0,2)
  d.TKI.prec<-1/(d.TKI.sd*d.TKI.sd)</pre>
  mu[2] ~ dnorm(0,.0001)
                                                # vague priors for trial baselines
  mu[5] ~ dnorm(0,.0001)
  mu[6] ~ dnorm(0,.0001)
  mu[7] ~ dnorm(0,.0001)
  mu[10] ~ dnorm(0,.0001)
  mu[11] ~ dnorm(0,.0001)
  MU ~ dnorm(0,.01)
                                               # prior for logit probabity with GEF
  MU.sd ~ dunif(0,2)
  MU.prec<-1/(MU.sd*MU.sd)
#Output
  # pairwise ORs and LORs for all possible pair-wise comparisons
  for (c in 1:(nt-1)) {
    for (k in (c+1):nt) {
      LOR[c,k] \leftarrow d[k] - d[c]
      OR[c,k] <- exp(LOR[c,k])</pre>
    }
  }
  # probability of AE
  for (k in 1:nt){
  logit(T[k])<-MU+d[k]</pre>
  }
```

#### G.1.2 Ast increase

}

```
model{
    # Binomial likelihood, logit link
```

```
# Model for relative treatment effects
  for(i in 1:ns){
                                   # LOOP THROUGH STUDIES
    for (k in 1:na[i]) {
                                       # LOOP THROUGH ARMS
      r[i,k] ~ dbin(p[i,k],n[i,k]) # binomial likelihood
      logit(p[i,k]) \leftarrow mu[i] + d[t[i,k]] - d[t[i,1]] # model for linear predictor
    }
  }
# Exchangeable relative treatment effects for TKIs
  for (k in 2:4){ d[k] ~ dnorm(d.TKI,d.TKI.prec) }
# Random effects model for absolute effects with GEF
  mu[1] ~ dnorm(MU,MU.prec)
  mu[3] ~ dnorm(MU,MU.prec)
  mu[6] ~ dnorm(MU,MU.prec)
  mu[7] ~ dnorm(MU,MU.prec)
# Priors
  d[1]<-0
                                               # treatment effect is zero for reference treatm
  for (k in 5:nt){ d[k] ~ dnorm(0,.001) }
                                               # vague priors for treatment effects
  d.TKI ~ dnorm(0,.001)
  d.TKI.sd ~ dunif(0,2)
  d.TKI.prec<-1/(d.TKI.sd*d.TKI.sd)</pre>
  mu[2] ~ dnorm(0,.0001)
                                             # vague priors for trial baselines
  mu[4] ~ dnorm(0,.0001)
  mu[5] ~ dnorm(0,.0001)
  mu[8] ~ dnorm(0,.0001)
  MU ~ dnorm(0,.01)
                                            # prior for logit probabity with GEF
  MU.sd ~ dunif(0,2)
  MU.prec<-1/(MU.sd*MU.sd)
```

```
for (k in (c+1):nt) {
                           LOR[c,k] \leftarrow d[k] - d[c]
                           OR[c,k] <- exp(LOR[c,k])</pre>
                   }
             }
             # probability of AE
             for (k in 1:nt){
             logit(T[k])<-MU+d[k]</pre>
}
G.1.3 Diarrhea
model{
       # Binomial likelihood, logit link
      # Model for relative treatment effects
             for(i in 1:(ns-1)){
                                                                                                                                      # LOOP THROUGH STUDIES
                    for (k in 1:na[i]) {
                                                                                                                                     # LOOP THROUGH ARMS
                           r[i,k] ~ dbin(p[i,k],n[i,k]) # binomial likelihood
                           logit(p[i,k]) \leftarrow mu[i] + d[t[i,k]] - d[t[i,1]] # model for linear predictor
                    }
             }
                    for (k in 1:na[ns]) {
                                                                                                                                        # LOOP THROUGH ARMS OF LAST STUDY (FLAURA)
                           r[ns,k] \sim dbin(p[ns,k],n[ns,k])
                                                                                                                                                    # binomial likelihood
                           logit(p[ns,k]) \leftarrow mu[ns] + d[t[ns,k]] - (0.66*d[1]+0.34*d[2]) # model for linear predictions and the second seco
                    }
      # Exchangeable relative treatment effects for TKIs
             for (k in 2:5){ d[k] ~ dnorm(d.TKI,d.TKI.prec) }
       # Random effects model for absolute effects with GEF
             mu[1] ~ dnorm(MU,MU.prec)
```

# pairwise ORs and LORs for all possible pair-wise comparisons

#Output

for (c in 1:(nt-1)) {

```
mu[3] ~ dnorm(MU,MU.prec)
  mu[5] ~ dnorm(MU,MU.prec)
  mu[7] ~ dnorm(MU,MU.prec)
  mu[10] ~ dnorm(MU,MU.prec)
  mu[11] ~ dnorm(MU,MU.prec)
  mu[13] ~ dnorm(MU,MU.prec)
  mu[14] ~ dnorm(MU,MU.prec)
# Priors
 d[1]<-0
                                                 # treatment effect is zero for reference treatm
  for (k in 6:nt) \{ d[k] \sim dnorm(0,.001) \}
                                                 # vague priors for treatment effects
 d.TKI ~ dnorm(0,.001)
 d.TKI.sd ~ dunif(0,2)
  d.TKI.prec<-1/(d.TKI.sd*d.TKI.sd)</pre>
 mu[2] ~ dnorm(0,.0001)
                                               # vague priors for trial baselines
 mu[4] ~ dnorm(0,.0001)
 mu[6] ~ dnorm(0,.0001)
 mu[8] ~ dnorm(0,.0001)
 mu[9] ~ dnorm(0,.0001)
 mu[12] ~ dnorm(0,.0001)
  mu[15] ~ dnorm(0,.0001)
 mu[16] ~ dnorm(0,.0001)
 MU ~ dnorm(0,.01)
                                              # prior for logit probabity with GEF
 MU.sd ~ dunif(0,2)
 MU.prec<-1/(MU.sd*MU.sd)
#Output
  # pairwise ORs and LORs for all possible pair-wise comparisons
  for (c in 1:(nt-1)) {
    for (k in (c+1):nt) {
      LOR[c,k] \leftarrow d[k] - d[c]
      OR[c,k] <- exp(LOR[c,k])</pre>
    }
```

```
}
    # probability of AE
    for (k in 1:nt){
    logit(T[k])<-MU+d[k]</pre>
    }
}
G.1.4 Dry skin
model{
  # Binomial likelihood, logit link
  # Model for relative treatment effects
    for(i in 1:(ns-1)){
                                        # LOOP THROUGH STUDIES
      for (k in 1:na[i]) {
                                        # LOOP THROUGH ARMS
        r[i,k] ~ dbin(p[i,k],n[i,k])
                                       # binomial likelihood
        logit(p[i,k]) \leftarrow mu[i] + d[t[i,k]] - d[t[i,1]] # model for linear predictor
      }
    }
      for (k in 1:na[ns]) {
                               # LOOP THROUGH ARMS OF LAST STUDY (FLAURA)
        r[ns,k] ~ dbin(p[ns,k],n[ns,k]) # binomial likelihood
        logit(p[ns,k]) \leftarrow mu[ns] + d[t[ns,k]] - (0.66*d[1]+0.34*d[2]) # model for linear predictions.
      }
  # Exchangeable relative treatment effects for TKIs
    for (k in 2:5){ d[k] ~ dnorm(d.TKI,d.TKI.prec) }
  # Random effects model for absolute effects with GEF
    mu[1] ~ dnorm(MU,MU.prec)
    mu[3] ~ dnorm(MU,MU.prec)
    mu[4] ~ dnorm(MU,MU.prec)
    mu[5] ~ dnorm(MU,MU.prec)
    mu[7] ~ dnorm(MU,MU.prec)
    mu[8] ~ dnorm(MU,MU.prec)
    mu[9] ~ dnorm(MU,MU.prec)
```

# Priors

```
d[1]<-0
                                                   # treatment effect is zero for reference treatm
    for (k in 6:nt) { d[k] ~ dnorm(0,.001) }
                                                   # vague priors for treatment effects
    d.TKI ~ dnorm(0,.001)
    d.TKI.sd ~ dunif(0,2)
    d.TKI.prec<-1/(d.TKI.sd*d.TKI.sd)</pre>
    mu[2] ~ dnorm(0,.0001)
                                                 # vague priors for trial baselines
    mu[6] ~ dnorm(0,.0001)
    mu[10] ~ dnorm(0,.0001)
    mu[11] ~ dnorm(0,.0001)
    MU ~ dnorm(0,.01)
                                                # prior for logit probabity with GEF
    MU.sd ~ dunif(0,2)
    MU.prec<-1/(MU.sd*MU.sd)
  #Output
    # pairwise ORs and LORs for all possible pair-wise comparisons
    for (c in 1:(nt-1)) {
      for (k in (c+1):nt) {
        LOR[c,k] \leftarrow d[k] - d[c]
        OR[c,k] <- exp(LOR[c,k])</pre>
      }
    # probability of AE
    for (k in 1:nt){
    logit(T[k])<-MU+d[k]</pre>
G.1.5 Eye problems
model{
  # Binomial likelihood, logit link
  # Model for relative treatment effects
```

```
# LOOP THROUGH STUDIES
 for(i in 1:ns){
   for (k in 1:na[i]) {
                                      # LOOP THROUGH ARMS
     r[i,k] ~ dbin(p[i,k],n[i,k]) # binomial likelihood
     logit(p[i,k]) \leftarrow mu[i] + d[t[i,k]] - d[t[i,1]] # model for linear predictor
   }
  }
# Exchangeable relative treatment effects for TKIs
  for (k in 2:nt){ d[k] ~ dnorm(d.TKI,d.TKI.prec) }
# Random effects model for absolute effects with GEF
 mu[1] ~ dnorm(MU,MU.prec)
 mu[2] ~ dnorm(MU,MU.prec)
# Priors
 d[1]<-0
                                               # treatment effect is zero for reference treatm
 d.TKI ~ dnorm(0,.001)
 d.TKI.sd ~ dunif(0,2)
 d.TKI.prec<-1/(d.TKI.sd*d.TKI.sd)</pre>
 MU ~ dnorm(0,.01)
                                            # prior for logit probabity with GEF
 MU.sd ~ dunif(0,2)
 MU.prec<-1/(MU.sd*MU.sd)
#Output
  # pairwise ORs and LORs for all possible pair-wise comparisons
 for (c in 1:(nt-1)) {
   for (k in (c+1):nt) {
     LOR[c,k] \leftarrow d[k] - d[c]
     OR[c,k] <- exp(LOR[c,k])</pre>
   }
  # probability of AE
  for (k in 1:nt){
```

```
}
}
G.1.6 Paronychia
model{
 # Binomial likelihood, logit link
 # Model for relative treatment effects
    for(i in 1:ns){
                                     # LOOP THROUGH STUDIES
     for (k in 1:na[i]) {
                                         # LOOP THROUGH ARMS
        r[i,k] ~ dbin(p[i,k],n[i,k])
                                       # binomial likelihood
        logit(p[i,k]) \leftarrow mu[i] + d[t[i,k]] - d[t[i,1]] # model for linear predictor
     }
    }
  # Exchangeable relative treatment effects for TKIs
    for (k in 2:4){ d[k] ~ dnorm(d.TKI,d.TKI.prec) }
  # Random effects model for absolute effects with GEF
   mu[1] ~ dnorm(MU,MU.prec)
   mu[3] ~ dnorm(MU,MU.prec)
   mu[4] ~ dnorm(MU,MU.prec)
   mu[7] ~ dnorm(MU,MU.prec)
   mu[9] ~ dnorm(MU,MU.prec)
  # Priors
   d[1]<-0
                                                 # treatment effect is zero for reference treatm
    for (k in 5:nt){ d[k] ~ dnorm(0,.001) }
                                                 # vague priors for treatment effects
   d.TKI ~ dnorm(0,.001)
   d.TKI.sd ~ dunif(0,2)
   d.TKI.prec<-1/(d.TKI.sd*d.TKI.sd)</pre>
   mu[2] ~ dnorm(0,.0001)
                                               # vague priors for trial baselines
```

logit(T[k])<-MU+d[k]</pre>

```
mu[5] ~ dnorm(0,.0001)
    mu[6] ~ dnorm(0,.0001)
    mu[8] ~ dnorm(0,.0001)
    mu[10] ~ dnorm(0,.0001)
    MU ~ dnorm(0,.01)
                                                # prior for logit probabity with GEF
    MU.sd ~ dunif(0,2)
    MU.prec<-1/(MU.sd*MU.sd)
  #Output
    # pairwise ORs and LORs for all possible pair-wise comparisons
    for (c in 1:(nt-1)) {
      for (k in (c+1):nt) {
        LOR[c,k] \leftarrow d[k] - d[c]
        OR[c,k] <- exp(LOR[c,k])</pre>
      }
    }
    # probability of AE
    for (k in 1:nt){
    logit(T[k])<-MU+d[k]</pre>
    }
G.1.7 Pneumonitis
model{
  # Binomial likelihood, logit link
  # Model for relative treatment effects
    for(i in 1:ns){
                                      # LOOP THROUGH STUDIES
      for (k in 1:na[i]) {
                                         # LOOP THROUGH ARMS
        r[i,k] ~ dbin(p[i,k],n[i,k])
                                        # binomial likelihood
        logit(p[i,k]) \leftarrow mu[i] + d[t[i,k]] - d[t[i,1]] # model for linear predictor
      }
    }
```

```
# Exchangeable relative treatment effects for TKIs
    for (k in 2:nt){ d[k] ~ dnorm(d.TKI,d.TKI.prec) }
  # Priors
    d[1]<-0
                                                   # treatment effect is zero for reference treatm
    d.TKI ~ dnorm(0,.001)
    d.TKI.sd ~ dunif(0,2)
    d.TKI.prec<-1/(d.TKI.sd*d.TKI.sd)</pre>
    mu[1] ~ dnorm(0,.01)
                                                   # priors for study effects
    mu[2] ~ dnorm(0,.01)
  #Output
    # pairwise ORs and LORs for all possible pair-wise comparisons
    for (c in 1:(nt-1)) {
      for (k in (c+1):nt) {
        LOR[c,k] \leftarrow d[k] - d[c]
        OR[c,k] <- exp(LOR[c,k])</pre>
      }
    }
    # probability of AE
    for (k in 1:nt){
    logit(T[k]) < -mu[2] + d[k]
}
G.1.8 Pruritus
model{
  # Binomial likelihood, logit link
  # Model for relative treatment effects
    for(i in 1:ns){
                                      # LOOP THROUGH STUDIES
      for (k in 1:na[i]) {
```

# LOOP THROUGH ARMS

```
r[i,k] ~ dbin(p[i,k],n[i,k]) # binomial likelihood
      logit(p[i,k]) \leftarrow mu[i] + d[t[i,k]] - d[t[i,1]] # model for linear predictor
    }
  }
# Exchangeable relative treatment effects for TKIs
  for (k in 2:5){ d[k] ~ dnorm(d.TKI,d.TKI.prec) }
# Random effects model for absolute effects with GEF
  mu[1] ~ dnorm(MU,MU.prec)
  mu[3] ~ dnorm(MU,MU.prec)
  mu[4] ~ dnorm(MU,MU.prec)
  mu[7] ~ dnorm(MU,MU.prec)
  mu[8] ~ dnorm(MU,MU.prec)
  mu[9] ~ dnorm(MU,MU.prec)
# Priors
  d[1]<-0
                                               # treatment effect is zero for reference treatm
  for (k in 6:nt){ d[k] ~ dnorm(0,.001) }
                                               # vague priors for treatment effects
  d.TKI ~ dnorm(0,.001)
  d.TKI.sd ~ dunif(0,2)
  d.TKI.prec<-1/(d.TKI.sd*d.TKI.sd)
  mu[2] ~ dnorm(0,.0001)
                                             # vague priors for trial baselines
  mu[5] ~ dnorm(0,.0001)
  mu[6] ~ dnorm(0,.0001)
  MU ~ dnorm(0,.01)
                                            # prior for logit probabity with GEF
  MU.sd ~ dunif(0,2)
  MU.prec<-1/(MU.sd*MU.sd)
#Output
  # pairwise ORs and LORs for all possible pair-wise comparisons
  for (c in 1:(nt-1)) {
```

```
for (k in (c+1):nt) {
                        LOR[c,k] \leftarrow d[k] - d[c]
                        OR[c,k] <- exp(LOR[c,k])</pre>
                  }
            }
            # probability of AE
            for (k in 1:nt){
            logit(T[k])<-MU+d[k]</pre>
}
G.1.9 Rash
model{
     # Binomial likelihood, logit link
      # Model for relative treatment effects
            for(i in 1:(ns-1)){
                                                                                                                             # LOOP THROUGH STUDIES
                  for (k in 1:na[i]) {
                                                                                                                            # LOOP THROUGH ARMS
                        r[i,k] ~ dbin(p[i,k],n[i,k]) # binomial likelihood
                        logit(p[i,k]) \leftarrow mu[i] + d[t[i,k]] - d[t[i,1]] # model for linear predictor
                  }
            }
                                                                                                           # LOOP THROUGH ARMS OF LAST STUDY (FLAURA)
                  for (k in 1:na[ns]) {
                        r[ns,k] ~ dbin(p[ns,k],n[ns,k]) # binomial likelihood
                        logit(p[ns,k]) \leftarrow mu[ns] + d[t[ns,k]] - (0.66*d[1]+0.34*d[2]) # model for linear predictions and the second seco
                  }
      # Exchangeable relative treatment effects for TKIs
            for (k in 2:5){ d[k] ~ dnorm(d.TKI,d.TKI.prec) }
      # Random effects model for absolute effects with GEF
            mu[1] ~ dnorm(MU,MU.prec)
            mu[3] ~ dnorm(MU,MU.prec)
            mu[6] ~ dnorm(MU,MU.prec)
            mu[8] ~ dnorm(MU,MU.prec)
```

mu[11] ~ dnorm(MU,MU.prec)

```
mu[12] ~ dnorm(MU,MU.prec)
 mu[14] ~ dnorm(MU,MU.prec)
  mu[15] ~ dnorm(MU,MU.prec)
# Priors
 d[1]<-0
                                                 # treatment effect is zero for reference treatm
 for (k in 6:nt){ d[k] ~ dnorm(0,.001) }
                                                 # vague priors for treatment effects
  d.TKI ~ dnorm(0,.001)
 d.TKI.sd ~ dunif(0,2)
  d.TKI.prec<-1/(d.TKI.sd*d.TKI.sd)</pre>
 mu[2] ~ dnorm(0,.0001)
                                               # vague priors for trial baselines
 mu[4] ~ dnorm(0,.0001)
 mu[5] ~ dnorm(0,.0001)
 mu[7] ~ dnorm(0,.0001)
 mu[9] ~ dnorm(0,.0001)
 mu[10] ~ dnorm(0,.0001)
 mu[13] ~ dnorm(0,.0001)
 mu[16] ~ dnorm(0,.0001)
  mu[17] ~ dnorm(0,.0001)
 MU ~ dnorm(0,.01)
                                              # prior for logit probabity with GEF
 MU.sd ~ dunif(0,2)
 MU.prec<-1/(MU.sd*MU.sd)
#Output
  # pairwise ORs and LORs for all possible pair-wise comparisons
 for (c in 1:(nt-1)) {
    for (k in (c+1):nt) {
      LOR[c,k] \leftarrow d[k] - d[c]
      OR[c,k] <- exp(LOR[c,k])</pre>
    }
  }
  # probability of AE
  for (k in 1:nt){
```

```
logit(T[k])<-MU+d[k]</pre>
           }
}
G.1.10 Stomatitis
model{
     # Binomial likelihood, logit link
     # Model for relative treatment effects
           for(i in 1:(ns-1)){
                                                                                                                       # LOOP THROUGH STUDIES
                 for (k in 1:na[i]) {
                                                                                                                   # LOOP THROUGH ARMS
                      r[i,k] ~ dbin(p[i,k],n[i,k])
                                                                                                                  # binomial likelihood
                      logit(p[i,k]) \leftarrow mu[i] + d[t[i,k]] - d[t[i,1]] # model for linear predictor
                 }
           }
                                                                                              # LOOP THROUGH ARMS OF LAST STUDY (FLAURA)
                 for (k in 1:na[ns]) {
                      r[ns,k] ~ dbin(p[ns,k],n[ns,k]) # binomial likelihood
                      logit(p[ns,k]) \leftarrow mu[ns] + d[t[ns,k]] - (0.66*d[1]+0.34*d[2]) # model for linear predictions and the second seco
                 }
     # Exchangeable relative treatment effects for TKIs
           for (k in 2:5){ d[k] ~ dnorm(d.TKI,d.TKI.prec) }
     # Random effects model for absolute effects with GEF
           mu[1] ~ dnorm(MU,MU.prec)
           mu[3] ~ dnorm(MU,MU.prec)
           mu[5] ~ dnorm(MU,MU.prec)
           mu[8] ~ dnorm(MU,MU.prec)
           mu[10] ~ dnorm(MU,MU.prec)
           mu[11] ~ dnorm(MU,MU.prec)
     # Priors
                                                                                                                                              # treatment effect is zero for reference treatm
           d[1] < -0
           for (k in 6:nt) { d[k] ~ dnorm(0,.001) }
                                                                                                                                              # vague priors for treatment effects
           d.TKI ~ dnorm(0,.001)
```

```
d.TKI.sd ~ dunif(0,2)
  d.TKI.prec<-1/(d.TKI.sd*d.TKI.sd)
 mu[2] ~ dnorm(0,.0001)
                                               # vague priors for trial baselines
 mu[4] ~ dnorm(0,.0001)
 mu[6] ~ dnorm(0,.0001)
  mu[7] ~ dnorm(0,.0001)
  mu[9] ~ dnorm(0,.0001)
  mu[12] ~ dnorm(0,.0001)
  mu[13] ~ dnorm(0,.0001)
 MU ~ dnorm(0,.01)
                                              # prior for logit probabity with GEF
 MU.sd ~ dunif(0,2)
  MU.prec<-1/(MU.sd*MU.sd)
#Output
  # pairwise ORs and LORs for all possible pair-wise comparisons
  for (c in 1:(nt-1)) {
    for (k in (c+1):nt) {
      LOR[c,k] \leftarrow d[k] - d[c]
      OR[c,k] <- exp(LOR[c,k])</pre>
    }
  }
  # probability of AE
  for (k in 1:nt){
  logit(T[k])<-MU+d[k]</pre>
  }
```

## G.2 Model parameter estimates

# H Value of hope

}

The value of hope measures the value patients place on tail-of-the-curve survival gains above and beyond what they place on increases in expected survival. It is computed for a given treatment strategy k relative to a reference treatment 1 by first estimating the certainty equivalent, or the life-years a patient would need to obtain to be indifference between treatment k and the reference treatment (treatment 1). To facilitate CEA, we work with QALYs rather than life-years. We assume a constant relative risk aversion utility function for QALYs x of the form,

$$u(x) = x^{\eta},\tag{A1}$$

where  $\eta$  is a measure of risks aversion with  $\eta > 1$  implying that a patient is risk loving and  $\eta < 1$  that a patient is risk averse. Using survey data from patients, Shafrin et al. (2017) estimate  $\eta = 1.39$  for NSCLC patients. The certainty equivalent,  $\alpha_{k1}$ , for treatment k relative to the reference treatment is computed by solving,

$$\int u(x + \alpha_{k1}) f_k(x) dx = V_1, \tag{A2}$$

where  $f_k(x)$  is the distribution of QALYs for treatment k and  $V_1 = \int u(x) f_1(x) dx$  is expected utility for the reference treatment. In the model,  $f_k(x)$  and  $f_1(x)$  are estimated using the distribution of QALYs from the individual-level simulation. We solve for  $\alpha_{k1}$  using the R root finding algorithm uniroot.

The value of hope,  $\theta$ , is the difference between the certainty equivalent and differences in expected survival,

$$\theta = \alpha_{k1} - [E_k(x) - E_1(x)],$$
(A3)

where  $E_k(x)$  and  $E_1(x)$  are mean QALYs for treatment k and the reference arm, respectively.

Note that since the algorithm is computationally intensive, we do not currently use the PSA to compute a distribution for  $\theta$ . Instead, we estimate a point estimate using the distribution of QALYs across all PSA iterations. Future work could speed up the algorithm so that it is fast enough for PSA by rewriting it in a compiled language such as C++.

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