MODELING AND SIMULATION APPLICATIONS WITH POTENTIAL IMPACT IN DRUG DEVELOPMENT AND PATIENT CARE

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DEDICATION

This work is dedicated to my thesis mentor, Dr. Robert R. Bies and my friends and family, especially my husband, David Yan, my upcoming baby, Austin, my parents, Michael and Maggie Li, my sister, Elsye, my grandparents, Ren-Jen and Shi-Hou Li, my parents-in-law, Zhaowan Yan and Jing Zhang, and my uncle and aunt, Li-Kang Wang and Ivy Li.
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Claire Li

Modeling and simulation applications with potential impact in drug development and patient care

Model-based drug development has become an essential element to potentially make drug development more productive by assessing the data using mathematical and statistical approaches to construct and utilize models to increase the understanding of the drug and disease. The modeling and simulation approach not only quantifies the exposure-response relationship, and the level of variability, but also identifies the potential contributors to the variability. I hypothesized that the modeling and simulation approach can: 1) leverage our understanding of pharmacokinetic-pharmacodynamic (PK-PD) relationship from pre-clinical system to human; 2) quantitatively capture the drug impact on patients; 3) evaluate clinical trial designs; and 4) identify potential contributors to drug toxicity and efficacy. The major findings for these studies included: 1) a translational PK modeling approach that predicted clozapine and norclozapine central nervous system exposures in humans relating these exposures to receptor binding kinetics at multiple receptors; 2) a population pharmacokinetic analysis of a study of sertraline in depressed elderly patients with Alzheimer’s disease that identified site specific differences in drug exposure contributing to the overall variability in sertraline exposure; 3) the utility of a longitudinal tumor dynamic model developed by the Food and Drug Administration for predicting survival in non-small cell lung cancer patients, including an exploration of the limitations of this approach; 4) a Monte Carlo clinical trial simulation approach that was used to evaluate a pre-defined oncology trial with a sparse drug concentration sampling schedule with the aim to quantify how well individual drug
exposures, random variability, and the food effects of abiraterone and nilotinib were determined under these conditions; 5) a time to event analysis that facilitated the identification of candidate genes including polymorphisms associated with vincristine-induced neuropathy from several association analyses in childhood acute lymphoblastic leukemia (ALL) patients; and 6) a LASSO penalized regression model that predicted vincristine-induced neuropathy and relapse in ALL patients and provided the basis for a risk assessment of the population. Overall, results from this dissertation provide an improved understanding of treatment effect in patients with an assessment of PK/PD combined and with a risk evaluation of drug toxicity and efficacy.

Robert R Bies, Pharm.D., Ph.D.
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<tr>
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<th>Description</th>
</tr>
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<tbody>
<tr>
<td>-2LL</td>
<td>-2 x log likelihood</td>
</tr>
<tr>
<td>AD</td>
<td>Alzheimer's disease</td>
</tr>
<tr>
<td>ADD</td>
<td>Additive</td>
</tr>
<tr>
<td>ALL</td>
<td>Acute lymphoblastic leukemia</td>
</tr>
<tr>
<td>ASW</td>
<td>African ancestry in Southwest USA</td>
</tr>
<tr>
<td>AUC</td>
<td>Area under the curve</td>
</tr>
<tr>
<td>BAV</td>
<td>Between-animal variability</td>
</tr>
<tr>
<td>BBB</td>
<td>Blood brain barrier</td>
</tr>
<tr>
<td>BQL</td>
<td>Below the quantitation limit</td>
</tr>
<tr>
<td>CA-125</td>
<td>Cancer antigen-125</td>
</tr>
<tr>
<td>CALGB</td>
<td>Cancer and Leukemia Group B</td>
</tr>
<tr>
<td>CHB</td>
<td>Han Chinese in Beijing</td>
</tr>
<tr>
<td>CL</td>
<td>Clearance</td>
</tr>
<tr>
<td>CNS</td>
<td>Central nervous system</td>
</tr>
<tr>
<td>COG</td>
<td>Children’s Oncology Group</td>
</tr>
<tr>
<td>CR</td>
<td>Complete remission</td>
</tr>
<tr>
<td>CRFS</td>
<td>Case report forms</td>
</tr>
<tr>
<td>CSF</td>
<td>Cerebrospinal fluid</td>
</tr>
<tr>
<td>CT</td>
<td>Computed tomography</td>
</tr>
<tr>
<td>CTCAE</td>
<td>Common terminology criteria of adverse event</td>
</tr>
<tr>
<td>CV</td>
<td>Coefficient of variation</td>
</tr>
<tr>
<td>CYP</td>
<td>Cytochrome P450</td>
</tr>
<tr>
<td>DOM</td>
<td>Dominant</td>
</tr>
<tr>
<td>ECF</td>
<td>Extracellular fluid</td>
</tr>
<tr>
<td>EQTL</td>
<td>Expression quantitative trait locus</td>
</tr>
<tr>
<td>FDA</td>
<td>Food and drug administration</td>
</tr>
<tr>
<td>FOCE</td>
<td>First order condition estimation</td>
</tr>
<tr>
<td>GEO</td>
<td>Gene expression omnibus</td>
</tr>
<tr>
<td>GTEX</td>
<td>Genotype-Tissue Expression</td>
</tr>
<tr>
<td>GWAS</td>
<td>Genome wide association study</td>
</tr>
<tr>
<td>IIV</td>
<td>Inter-individual variability</td>
</tr>
<tr>
<td>KA</td>
<td>Absorption constant</td>
</tr>
<tr>
<td>KTR</td>
<td>Transit constant</td>
</tr>
<tr>
<td>LASSO</td>
<td>Least absolute shrinkage and selection operator</td>
</tr>
<tr>
<td>LLOQ</td>
<td>Lower limit of quantitation</td>
</tr>
<tr>
<td>MAF</td>
<td>Minor allele frequency</td>
</tr>
<tr>
<td>MBDD</td>
<td>Model based drug development</td>
</tr>
<tr>
<td>NA</td>
<td>No available</td>
</tr>
<tr>
<td>NC</td>
<td>No change</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
</tr>
<tr>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>NCI</td>
<td>National cancer institute</td>
</tr>
<tr>
<td>NE</td>
<td>Neuropathy</td>
</tr>
<tr>
<td>NEP</td>
<td>Neuropathy+neuropathic pain</td>
</tr>
<tr>
<td>NNE</td>
<td>Nearest neighbor estimation</td>
</tr>
<tr>
<td>OFV</td>
<td>Objective function value</td>
</tr>
<tr>
<td>PD</td>
<td>Pharmacodynamics</td>
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<tr>
<td>PET</td>
<td>Positron emission tomography</td>
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<td>PGP</td>
<td>P-glycoprotein</td>
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<tr>
<td>PK</td>
<td>Pharmacokinetics</td>
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<td>PM</td>
<td>Pharmacometrics</td>
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<td>POP-PK</td>
<td>Population pharmacokinetic</td>
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<td>PSA</td>
<td>Prostate specific antigen</td>
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<tr>
<td>Q</td>
<td>Intercompartmental clearance</td>
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<tr>
<td>REC</td>
<td>Recessive</td>
</tr>
<tr>
<td>RECIST</td>
<td>Response evaluation criteria in solid tumor</td>
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<tr>
<td>ROC</td>
<td>Receiver operating characteristics</td>
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<tr>
<td>SAEM</td>
<td>Stochastic approximation expectation maximization</td>
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<tr>
<td>SC</td>
<td>Subcutaneously</td>
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<tr>
<td>SLD</td>
<td>Sum of longest diameter</td>
</tr>
<tr>
<td>SNP</td>
<td>Single nucleotide polymorphism</td>
</tr>
<tr>
<td>TRS</td>
<td>Treatment resistant schizophrenia</td>
</tr>
<tr>
<td>V</td>
<td>Volume of distribution</td>
</tr>
<tr>
<td>VIPN</td>
<td>Vincristine-induced peripheral neuropathy</td>
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</table>
CHAPTER I: Introduction

1. The need for modeling and simulation in drug development

Despite significant expenditures, the drug development process requires a substantial amount of time and it is expensive and prone to failures. The average length of clinical development is quoted as 7-12 years at a cost of $0.8-1.7 billion per approved agent (Kaitin et al., 2010; Dimasi et al., 2001; Connolly et al., 2001; Lesko et al., 2004). The cost of drug development takes into account those compounds that undergo clinical testing but do not make it to the marketing approval or fail NDA approval. Therefore, the rate at which pharmaceutical firms successfully develop investigational compounds for marketing approval by regulatory agencies is a critical indicator of the effectiveness of the drug development process (DiMasi et al., 2010). A retrospective analysis of both public and private drug pipeline database as well as from surveys comprising nearly 4000 drugs and biologics from the 50 largest pharmaceutical firms between the mid-1990s and the early 2000s by DiMasi’s group pointed to clinical approval success rates that remained lower than 20% with an upward trend in costs (DiMasi et al., 2010). Estimated clinical approval success rates differed significantly by therapeutic class; for example, in oncology, the success rate was only ~5% (Kola et al., 2004). Furthermore, even with the massive amount of data that are generated and obtained from the clinical trials every year, lots of useful knowledge and information remains undiscovered or underutilized (Ette et al., 2007). Processes and technological innovations that can improve the predictability of outcomes for new compounds can therefore significantly increase the productivity of new drug innovation (DiMasi et al., 2010). Indeed, in the white paper Challenge and Opportunity on the Critical Path to New Products published in March of 2004, the Food
and Drug Administration (FDA) has addressed this concern and states “Not enough applied scientific work has been done to create new tools to get fundamentally better answers about how the safety and effectiveness of new product that can be demonstrated in faster time frames, with more certainty, and at lower cost. A new product development toolkit—containing powerful new scientific and technical methods such as animal or computer-based predictive models, biomarkers for safety and effectiveness, and new clinical evaluation techniques—is urgently needed to improve predictability and efficiency along the critical path from laboratory concept to commercial product. We need superior product development science to address these challenges.” To highlight all these issues regarding the high attrition rates and increasing costs in drug development, model-based drug development has been proposed as one methodology to potentially make drug development more productive by assessing the ongoing clinical trial studies as well as historical drug databases using mathematical and statistical approaches to construct and utilize models to increase the understanding of the drug and disease.

2. Model based drug development (pharmacometrics and statistics)

Model-based drug development (MBDD) is the concept of utilizing pharmacostatistical approaches to evaluate drug efficacy and safety from preclinical through the clinical data to improve drug development knowledge management and decision-making (Milligan et al., 2013; Lalonde et al., 2007). As Lalonde et al. described, the key components of MBDD can be divided into six categories: PK-PD and disease/placebo model; meta-analysis of candidate drug and competitor data, design considerations and trial execution models; data-analytic models; quantitative decision criteria; and trial performance metrics (Milligan et al., 2013; Lalonde et al., 2007). Those principles can
potentially overcome the common limitation of the current drug development strategy, specifically, that prior information is partly or completely ignored when analyzing and interpreting the results of the most recent clinical trials.

One of the essential parts of MBDD is the discipline of pharmacometrics. Pharmacometrics has been described as “the science of developing and applying mathematical and statistical methods to a) characterize, understand, and predict a drug’s pharmacokinetic and pharmacodynamic behavior, b) quantify uncertainty of information about that behavior and c) rationalize data-driven decision making in the drug development process and pharmacotherapy” (Ette et al., 2007). The modeling and simulation approach not only quantifies the exposure-response relationship, disease progression, and the level of variability, but also identifies the potential contributors to that variability. In addition, these models can be linked and applied to competing study designs and to customize patient drug therapy through therapeutic drug monitoring and improved population dosing strategies; therefore, facilitating the implementation of the personalized medicine (Bonate, 2011). The idea of personalized medicine can be further utilized to emphasize the significance of biomarker findings. As a result, biomarker identification and validation become one of the major foci of pharmacometricians across disciplines. Applying quantitative assessment approaches to understand biomarker dynamics can potentially: identify patients at risk for a disease; predict a patient response; and predict the risk of toxicity.

Other statistical tools have also become popular to describe the relationship of disease and clinical outcome which may or may not involve drugs or treatments using mathematical equations such as regression analysis and survival analysis. In particular,
survival analysis has been commonly used in clinical studies to analyze time to event data which can provide insight of patients’ efficacy and safety profiles and facilitate decision making such as dose selection or patient care.

Since the quantity of the data expands as the improvement of data collection method, an efficient computer tool becomes critical and demanded to facilitate the access to large amount of information (ex: pharmacogenomics profiles). This can be overcome by the development and implementation of bioinformatics tool.

3. The opportunities (application) of modeling and simulation in drug development

As the role of modeling and simulation in drug development becomes more significant, there are many opportunities available for modeling across different phase of studies. In the journal of the pharmaceutical sciences 2002, Meibohm and Derendorf initially indicated the potential modeling and simulation applications from the discovery and preclinical phase to clinical setting and even the post-marketing phase in Figure 1.1. As shown in Figure 1.1, in the early stages of drug development, modeling and simulation can explore the pharmacokinetic and pharmacodynamic of drug and drug-drug interaction in \textit{in vivo} system and leverage the understanding of biomarker to evaluate drug efficacy and toxicity profile. It can also be useful in selecting dosage regimen or form based on the previously developed PK/PD relationship and integrate the early decision making in candidate selection. Furthermore, with the modeling assessment of preclinical studies, a first to human dose can be suggested by extrapolating from the \textit{in vivo} model, and the translational approach can be also beneficial for systematically assessing some pharmacological relationships which have not been assessable in the human system. A parallel approach has been suggested by combining preclinical and
early clinical development as a way to expand the learning process to all the phases of drug development. In the learning and confirming process of clinical trial, modeling and simulation serves as a tool to continuously evaluate exposure-response relationship across different therapeutic areas (Huang et al., 2013; Meibohm et al., 2001; Sheiner, 1997). The model can further evaluate contributors such as gender and food effects on exposure, predict PK/PD under a new dosing regimen, predict PK/PD in a special population such as children or elderly, and characterize drug-drug interaction and drug-disease interaction (Meibohm et al., 2002). In clinical drug development, predictive tools such as Monte Carlo simulation can explore various dosage regimens and optimize the trial design which might allow reducing the numbers and costs of the studies and improving drug development efficiency (Meibohm et al., 2002).

Figure 1.1. Opportunities (applications) for modeling and simulation across the drug development process. (This figure is reproduced with permission of John Wiley & Sons, Inc. Meibohm B anad Derendorf H, Pharmcokinetic-pharmcodynamic studies in drug product

4. Type of models

As the demands and opportunities of modeling and simulation increases in each phase of drug development, various types of models have been advocated to address specific questions. Some of the model strategies are introduced in the following sections, and the applications are demonstrated in each section.

4.1 Pharmacokinetic modeling

Pharmacokinetics began as a way to characterize the absorption, distribution, metabolism and elimination of a drug in the body and to reduce a concentration-time profile into a set of parameters that could be used for comparison, evaluation, and prediction (Teorell, 1937; Teorell 1937; Widmark, 1933; Wanger, 1973; Widmark, 1933; Derendorf et al., 2000). As the assays or the measurement tools becoming available not only in the plasma but also in some tissue levels, disposition of drug exposure pattern can be better understood. Furthermore, many forms of drug exposure were also characterized in the pharmacokinetic analysis. Depending on whether drug concentration measurements are available; sometimes, dose can be used as a representation of nominal exposure, and population average exposures after dose have been proposed. These are known as KPD models explore the dose-driven shape of the concentration vs time profile specifically the temporality without inter-individual variability in the context of exposure and pharmacodynamics relationship. Typically, drug concentrations measured in plasma represent the systemic exposure of the drug. Pharmacokinetic parameters such as clearance (CL), volume of distribution (Vd) or descriptors such as area under the curve (AUC), maximum concentration (Cmax) and elimination half–life ($t_{1/2}$) can be derived.
or calculated. Beyond the drug concentrations measured in the biologic fluid, with the innovation and improvement of the measurement techniques, drug concentrations now can be measured at the target site using microdialysis or positron-emission tomography scan so that the responses can be more precisely correlated to drug concentrations at the effect site (Derendorf et al., 2000; Chefer et al., 2009; Jacobson et al., 2013). In addition, understanding the covariate effect on systematic exposure and variability associated with exposure is also an important component in a pharmacokinetic model. Covariates are characteristics describing the patients, the conditions of the drug treatment or other factors potentially influencing the outcome. In late 90’s, Vozeh’s and Grasela’s groups demonstrated how covariates can be used to predict patients’ pharmacokinetics through the covariate model development in a population analysis (Vozeh et al., 1982; Grasela et al., 1984; Grasela et al., 1985; Grasela et al., 1987). The covariates can be constant within an individual or time-varying. They not only contribute to the estimates of structure model parameters but also impact the random effects distribution (Ribbing, 2007). In general, covariates can be classified as intrinsic factors (e.g. age, race, weight, sex, and genotype) or extrinsic factors (e.g. compliance, and smoking status) or can be categorized as categorical or continuous variables based on the measurement scale (Holford, 2013). In the process of model development, an assessment of covariate effect can improve predictive model performance for subject in the current data set, for trial simulation of future studies or for future patient population (Gastonguay MR, 2011).

4.2. Pharmacodynamic modeling

Pharmacodynamics is often defined as what the drug does to the body (Ette et al., 2007). Based on the definition from Derendorf et al, “pharmacodynamics is a broad term
that is intended to include all of the pharmacological actions, pathophysiological effects and therapeutic responses, either beneficial or adverse of active drug ingredient, therapeutic moiety, and/or its metabolite(s) on the various systems of the body from subcellular effects to clinical outcomes” (Derendorf et al., 2000). In pharmacodynamic modeling, the pharmacological action is often modeled as a time dependent effect that is related to drug exposure, and the endpoints of pharmacodynamics can be physiological changes such as blood pressure and cholesterol level or clinical outcomes such as survival or toxicity. In addition, with the flexibility of mathematical functions, different types of pharmacodynamic endpoints including continuous, categorical, ordered categorical or count data can be described. Another type of pharmacodynamic response is the time to a specific event. A common approach for modeling, this type of event is survival analysis. Typically, a proportional hazard model or parametric survival model is likely to be used to describe the baseline hazard and the risk that depends on covariates in a predefined time period.

4.3. Population PK/PD modeling

Sheiner and his colleagues published several articles illustrating a new mathematical approach to analyze pharmacokinetic data, which was later called population pharmacokinetics in the late 1970s and early 1980s (Sheiner et al., 1977; Sheiner et al., 1980; Sheiner et al., 1981; Sheiner et al., 1982; Bonate, 2005). In one of the Sheiner’s publications, he demonstrated that less-biased estimates of the population means and variances were found using the population approach than either the naïve-pooled or 2-stage approach (Sheiner et al., 1981; Bonate, 2005). Population methods were further applied to assess pharmacodynamics endpoints. Population PK/PD modeling
approaches represent a methodology of leveraging sparse PK and PD data. Commonly, this approach is applied to relatively sparse sampling but in a sufficient number of individuals so that inter-individual variability can be assessed and allows integrating data from single or multiple studies as well as studies from different phases (Ette et al., 2007; Gross et al., 2000). With the flexibility of analyzing combined data from different studies, PK or PD profile of the population can be better represented. However, analyzing combined data in a typical model-based metal-analysis may potentially result in bias estimates of parameters. Therefore, this strategy should be carefully evaluated and applied (Gastonguay et al., 2005; Gastonguay et al., 1999). This strategy also allows for the identification and quantification of the sources of inter-individual variability in response such as genetic polymorphisms in metabolic enzyme or transporter, clinical and demographic factors. In particular, the population modeling approach has become one of the standard assessments of clinical drug development that regulatory agencies recommend (Department of Health and Human Services, 1999).

4.4. Biomarker modeling

The official National Institutes of Health (NIH) definition of a biological marker (Biomarker) is “a characteristic that is objectively measured and evaluated as an indicator of normal biological processes, pathogenic processes, or pharmacologic responses to a therapeutic intervention.” This definition is broad and though not explicitly stated, includes laboratory tests, radiologic studies as well as physical exam findings. Although biomarker is a relatively new term that dates back to the late 1960s, biologic assessments and measurements in the evaluation of human disease were practiced in antiquity (Ferguson et al., 2010). Clinically useful biomarkers have evolved over time and play an
increasingly important role in many aspects of pharmaceutical discovery and
development including personalized medicine and the assessment of safety profile,
reflecting the scientific and technologic progress made over the centuries.

As a result, an increasing number of clinically relevant tests and procedures are
available to estimate organ injury and guide treatment. Recent discoveries in genetics and
molecular biology have resulted in impressive advances in our understanding of the
pathophysiologic processes of individual disease and yielded an abundance of
prospective therapies directed against novel targets (Ferguson et al., 2010). This has
brought about an increased focus on biomarker identification, validation, and
quantification, as well as the development of analytical technologies for biomarker
measurement. Efforts at biomarker discovery and validation have intensified since the
twenty-first century (Ferguson et al., 2010). Advanced genomic, proteomic, and
metabolomics techniques now permit comparative analysis of specimens from healthy
and diseased individuals, facilitating biomarker identification. Generally, there are four
types of biomarker endpoints (Figure 1.2) (Jenkins et al., 2011). The first is called a
prognostic endpoint which is used to predict the likely disease prognosis independent of
the mode of treatment. For instance, the number of circulating tumor cells (CTCs) in
peripheral blood at baseline can give an indication of survival prognosis in prostate
cancer (De Bono et al., 2008). The second type is a predictive biomarker that predicts the
likelihood of response to a particular treatment or a class of treatments. As an illustration,
CTCs can also be a predictive biomarker when compared wild type patients vs. patients
with mutation in the same treatment. This acts more like a covariate to explain the
variation in the outcome. The third type of biomarker is a pharmacodynamics biomarker.
The definition of a pharmacodynamic biomarker is a response over time to a treatment intervention. Simple biomarker examples include cancer antigen 125 for ovarian cancer (Schmidt et al., 2011) and PSA for prostate cancer (Romero et al., 2013), and tracking treatment over time is an important component. Pharmacodynamic biomarkers could reflect the safety and/or efficacy of a treatment and are typically measured at multiple time points. Markers that correlate well with a widely accepted clinical outcome at both an individual and group level could potentially act as a surrogate endpoint and substitute for a recognized clinical endpoint (Katz et al., 2004; Rothmann et al., 2012) and are the fourth type of biomarker. For example, LDL cholesterol can act as a surrogate for major cardiovascular events in the licensing of statins (Tardif et al., 2006). The model based assessment of the dynamics of a biomarker can provide insight into the disease progression and capture a therapeutic drug effect which might be useful for early decision making before the clinical endpoints are available.
Figure 1.2. **Biomarker endpoint types** (This figure is reproduced with permission of John Wiley & Sons, Inc. Jenkins et al., A statistician’s perspective on biomarkers in drug development, and copyright © 2011 John Wiley & Sons, Ltd.).

4.5. Clinical Trial simulation

Another significant component of pharmacometrics is stochastic simulation. Simulation has been widely used in many disciplines such as engineering before being used in drug development. It can serve not only as a predictive tool to evaluate the proposed trial design and the outcome under certain assumptions and to exposure pattern resulting from various dosage regimens, sampling schemas and population characters but also as an analytical tool to characterize exposure-response relationship in different modeling circumstances in a virtual population (Kimko et al., 2002; Kimko et al., 2011; Meibohm et al., 2002). Clinical trial simulation also allows us to compare different study designs and optimize study design by updating the design continuously. With the simulation assessment, the drug development process can be more efficient and powerful and more informative in bridging studies from one phase to another.

4.6. Time to event model (GWAS study)

Clinical trials are often performed to evaluate the efficacy of new treatment regimens and the efficacy of existing treatments in a special population (National cancer institute, 2011). As mentioned briefly in section 4.2, the efficacy endpoints of the clinical trials are primary described as an event of interest such as death, relapse, and adverse drug reaction (Crom et al., 1994). The follow-up time of the patients may vary depending on the disease and treatment, the time component is an important assessment to understand the event occurrence. These statistical models consider the time course until an event occurs and compare the cumulative probability of events over time for two or
more cohorts (Lee et al., 1982; Singh et al., 2011). Therefore, a statistical procedure which is likely to applied for analyzing this endpoints is called time to event analysis. Because survival is one of the most common endpoints in time to event analysis, people also describe this type of analysis as survival analysis. In time to event analysis, one of the essential components is how the dichotomous event of interest is pre-defined. For example, if an adverse drug reaction is classified using a grading scale from 0 to 5, the event of interest can be specified as binary or categorical outcome based on the question we are interested in (National cancer institute, 2009). Another key component of most time to event model is censoring. Censoring can arise under three circumstances: when an individual does not experience the event during the study period; when an individual discontinues the follow-up during the study period; and when an individual drops out from the study due to other events (Singh et al., 2011).

One of the most common time to event model is the Cox proportional hazards model (Cox et al., 1972). This model was first introduced by Cox, in 1972, for analysis of survival data with and without censoring, and for identifying the contributors to the difference of survival in clinical trials. These contributors include treatment effect, prognostic and clinical covariates. Different from parametric survival or the Kaplan-Meier method (Itman et al., 1992), the Cox regression model is considered a semi-parametric method which does not require a pre-specified hazard function for the baseline hazard but considers the hazard to be proportional at any point in time for the characteristics based on the model assumption. The hazard is calculated using the equation shown below:

\[ H(t) = H_0(t) \exp(b_1X_1 + b_2X_2 + b_3X_3 + \cdots + b_jX_j) \]
where $X_1 \ldots X_j$ represent a group of predictor variables, and $H_0(t)$ is the baseline hazard at time $t$, representing the hazard for a person when all the predictor variables are 0 (Cox et al., 1972).

The Cox model has been widely used in covariate selection for association studies such as Genome wide association study (GWAS), and it is very useful and robust tool in clinical research and provides valuable information about an intervention.

5. Hypothesis and specific aims

Modeling and simulation has been shown to be a critical tool in drug development and patient care. In drug development, modeling and simulation provides an early assessment of a new drug entity in the initial discovery phase to facilitate: the compound selection; the prediction of pharmacokinetic-pharmacodynamic (PK-PD) relationships from in-vivo animal results to human; the characterization of exposure-response profiles at both the individual and population levels in the clinical studies; the evaluation of the trial design; and insights to inform decision making in drug development. Furthermore, as the studies go beyond drug approval, modeling and simulation can also be applied to the post-marketing evaluations for special populations and to predict drug toxicity and efficacy in a long-term patient care setting with the potential to inform clinical recommendations.

The main objectives of this proposal use a modeling and simulation approach: 1) to leverage our understanding of PK/PD relationship from pre-clinical system to human; 2) to quantitatively understand the drug impact on patients; 3) to evaluate clinical trial designs and; 4) to identify potential contributors (predictors) to drug toxicity and efficacy. This work explores the hypothesis that modeling and simulation can be useful to provide
insight into trial designs and facilitate decision making (Steimer et al., 2000). To test this hypothesis, we proposed the following specific aims:

1. Build a population pharmacokinetic model that accounts for both plasma and brain concentrations measured in rats and utilize this model to predict human concentrations of clozapine and its N-desmethyl metabolite, norclozapine. (Preclinical PK model)

2. Identify covariates that contribute to variability in sertraline concentration by performing a population pharmacokinetic analysis of sertraline in elderly patients with Alzheimer disease and generating population pharmacokinetic parameters for this population. (Population PK model in a special population)

3. Assess and refine the published FDA longitudinal tumor size model for predicting survival in NSCLC patients using archived tumor measurement data. (Pharmacodynamic/Biomarker model)

4. Evaluate whether a pre-defined oncology trial with a sparse drug concentration sampling schedule can adequately capture individual level drug exposures, random variability and the food effects of abiraterone and nilotinib. (Clinical trial simulation and trial design evaluation)

5. Establish a candidate gene database which included those polymorphisms associated with vincristine-induced neurotoxicity in childhood ALL patients from the GWAS study and gene enrichment analysis (Time to event model)

6. Create a signature comprising a combination of genetic and clinical markers that predict vincristine-induced neurotoxicity and relapse in childhood ALL patients using LASSO penalized regression model. (Signature model)
Successful completion of these aims would result in accelerated and information rich decision making by leveraging data from previous trials to simulate clinical outcomes as well as aid in the design of future clinical trials. Modeling and simulation allows one to explore the pharmacokinetic profile in a special population and exposure and response relationship in preclinical and clinical setting to facilitate drug development as well as optimal drug utilization and therefore provide better care for the patients. Furthermore, potential biomarkers for survival prediction and drug-induced adverse reactions identified by the modeling approach can help in guiding treatment decisions and facilitating the future protection for patients. Ideally, patients most likely to derive a benefit from treatment are more likely to be targeted; in contrast, those patients identified as non-responders are more likely to be excluded so that unnecessary harm to patients can be minimized. This may allow for the identification of ineffective treatments earlier in the clinical trial process thus also avoiding unnecessary patient exposures. In this study, modeling and simulation demonstrates the potential applications in drug development and patient care and the utility of clinical prediction.
CHAPTER II: Preclinical model-prediction of brain clozapine and norclozapine concentrations in humans using a scaled pharmacokinetic model for rat brain and plasma pharmacokinetics (Published in Journal of translational medicine with open access license http://www.biomedcentral.com/about/license) (This material is reproduced with permission of BioMed Central)

1. Introduction

Schizophrenia is a debilitating disorder that affects approximately 1% of the global population without regard to race, sex or socioeconomic status (Mathers et al., 2006). It typically strikes in the late-teens or early twenties and is characterized by a high rate of morbidity and mortality. Given these high personal and societal costs, investment in research aimed at understanding the biology of the disease, its genetic components and their interplay with environmental factors, continues on many levels. Over the past 50 years, pharmacotherapeutic support has been instrumental in managing primarily the positive symptoms of the disease and hinges on suppression of a central circuitry dysfunction that can be normalized by antagonism of dopamine D2 receptors in the striatum (Murray et al., 2008). Introduction of clozapine, the first so-called atypical antipsychotic approximately 25 years ago, represented a significant advance in our understanding of schizophrenia from a systems biology perspective in that this drug did not have the typical side effects of the 1st generation neuroleptics. This reduction in side effects was attributed to higher 5HT-2A than D2 binding (Meltzer et al., 1989).

However, clozapine pharmacology is not limited to D2 and 5HT2A antagonism. Albeit unintentionally, the drug binds to several other dopamine and serotonin receptor subtypes, as well as to muscarinic M1/M4 and alpha-1 adrenergic receptors with
pharmacologically relevant affinity (Horacek et al., 2006). From a clinical perspective, this broad receptor coverage may account for clozapine’s unique superiority in treatment resistant schizophrenia (TRS), even amongst other atypical antipsychotics. From a research perspective, the broad receptor coverage of clozapine conceivably makes the drug a useful tool to advance our understanding of complex pharmacotherapy that incorporates multiple interacting receptor systems.

The use of positron emission tomography (PET) imaging to measure receptor occupancy of clozapine and other atypical antipsychotics in humans has been invaluable in demonstrating the importance of D2 and 5HT2A receptor antagonism contributing to the efficacy of these drugs (Takeuchi et al., 2013; Moriguchi et al., 2013; Tsuboi et al., 2013). However, broader application of this non-invasive technique has been limited by the lack of ligands specific for other receptors to which clozapine has affinity. In this regard, availability of other approaches that are complementary to PET imaging would be useful. One possibility is to link non-clinical measurements of clozapine disposition in the brain with clinical studies of clozapine systemic exposure using a translational PK modeling approach. Prediction of clozapine CNS exposure could then be related to its receptor binding kinetics at multiple receptors to impart a virtual predicted pharmacodynamic component to a model. This approach has been used recently to predict CNS concentrations of atomoxetine and duloxetine that were in the range of receptor affinities associated with therapeutic doses (Kielbasa et al., 2012). In a related manner, a population pharmacokinetic-pharmacodynamic (PK-PD) modeling approach was used to predict D2 receptor occupancy of olanzapine in humans (Johnson et al., 2011), and the D2 and 5HT2A receptor occupancy of risperidone and its active
metabolite paliperidone (9-OH risperidone) (Takeuchi et al., 2013; Moriguchi et al., 2013; Tsuboi et al., 2013; Kozleiska et al., 2012). These studies, as well as earlier PK-PD models applied to other CNS drugs (de Lange et al., 2012; de Lange et al., 2005), provide confidence in the ability of this approach to deepen our understanding of drug action in human brain.

A recent study measured clozapine and its N-desmethyl metabolite, norclozapine, in extracellular fluid (ECF) of rat medial prefrontal cortex using quantitative microdialysis, and these results provided evidence of net efflux from brain across the blood-brain barrier (BBB) (Cremers et al., 2012). This suggests that plasma concentrations may not be a good predictor of brain concentration for clozapine or norclozapine. Therefore, prediction of clozapine exposure in the ECF of human brain using a translational PK modeling approach could be cross-validated against PET results at D2 and 5HT2A receptor occupancy in humans, and subsequently used to estimate clozapine receptor occupancy at the drug’s other receptor targets for which PET tracers do not exist. Such comprehensive PK-PD model could potentially support individualized dosing of clozapine to improve its efficacy and CNS tolerability. It would also support research aimed at discovering new approaches for the treatment of the schizophrenia in its different forms.

The purpose of this study was; (1) to build a PK model that accounted for both plasma and brain concentrations measured in rats; (2) to utilize this model to predict concentrations of clozapine and norclozapine in human brain. This would allow for the prediction of expected receptor occupancy in humans.
2. Materials and Methods

2.1. Study Design

A single dose of clozapine (10mg/kg) was administered subcutaneously to four male Wistar rats with an average weight of 0.35 kg purchased from Harlan (Zeist, The Netherlands). Three days prior to administration a microdialysis guide cannula was surgically implanted in the medial prefrontal cortex; at the same time, a catheter for blood sample collection was placed in the right jugular vein and was exteriorized through an incision at the top of the head. This vascular cannulation enabled an equivalent volume of saline replacement for each blood sample. A MetaQuant probe (6 mm, cellulose membrane, BrainLink, The Netherlands) was inserted into the guide cannula 24 hours prior to drug administration to enable sampling of brain extracellular fluid (ECF). Concentrations of clozapine and norclozapine were measured in plasma and brain (ECF) by HPLC with tandem mass spectrometry in the positive ion mode as previously described (Cremers et al., 2012). For each rat, the unbound concentrations in each compartment were measured at 9 time points (0, 15, 30, 60, 90, 120, 240, 360 and 480 minutes) in plasma and 18 time points (-30, 0, 30, 60, 90, 120, 150, 180, 210, 240, 270, 300, 330, 360, 390, 420, 450 and 480 minutes) in brain. A single dose of norclozapine (10 mg/kg) was also administered subcutaneously to another five male Wistar rats with an average weight of 0.34 kg bought from the same Harlan laboratories. Concentrations were measured in plasma and brain ECF, and the same time points were used as those specified for clozapine.
2.2. Model Development

Different model structures were initially evaluated using the system dynamics software VENSIM (Ventana Systems, Inc., MA, US). Thereafter a population approach was used to describe the pharmacokinetics of clozapine and norclozapine. Population PK parameters were estimated using a nonlinear mixed effect modeling approach, as implemented in NONMEM version 7.2 (Icon Development Solutions, Hanover, Maryland) using Wings for NONMEM version 7 (Holford, 2013). The first-order conditional estimation method (FOCE) with interaction was used to estimate the structural PK parameters and the random effects parameters.

Model development was started with an assessment of clozapine PK in plasma. One and two compartment models with first order absorption for clozapine in plasma were tested. A peripheral compartment structure was subsequently implemented to represent the brain extracellular fluid concentrations. The transfer characteristics of clozapine between the plasma and the brain compartment were evaluated using an intercompartmental clearance, $CL_{in}/CL_{out}$, as well as incorporating delay functions (Savic et al., 2007). These delay functions included a lag time and transit compartment approaches. Once the structural model for clozapine was established, the plasma compartment of norclozapine was integrated and then connected to the brain compartment. The same strategy was utilized in building the structural model for norclozapine concentrations that were measured following norclozapine administration. Clozapine and norclozapine concentration measurements were then combined from the 9 rats and modeled simultaneously in the final structural model. The volume of distribution of clozapine and norclozapine in brain were tested with and without fixing this parameter.
to a literature reported value (Fridén et al., 2007). A parallel metabolic pathway from the extravascular space was also explored.

Between-animal variability (BAV) for PK parameters was assumed to be log-normally distributed and evaluated using an exponential model $P_i = P_{TV} \times e^{\eta_p}$ where $P_i$ is the parameter estimate for the $i^{th}$ animal, and $P_{TV}$ is the typical parameter value at the population level. The difference between $i^{th}$ individual and population parameter values was described by $\eta_p$, which was identically distributed with mean equal to 0 and variance, $\omega_{\eta}^2$ (Feng et al., 2006). A combined additive and proportional model was first used to describe the intra-animal variability. If one of the elements of the model was found to be negligible and not significant, it was then removed from the residual error model.

Residual error parameters were assumed to be normally distributed with mean equal to 0 and variance, $\sigma^2$.

2.3 Model selection and evaluation

Model evaluation was based on a likelihood ratio test using the objective function value (OFV) provided by NONMEM. The minimum OFV returned by NONMEM is approximately equal to $-2 \times \log$ likelihood ($-2LL$) and served as a guide during model design. A decrease in $-2LL$ of 6.63 points for 1 degree of freedom was regarded as a significant model improvement, corresponding to a p value of 0.01 for nested models.

The final model was further examined using goodness-of-fit plots generated using R version 2.13 based on the conditional weighted residuals distribution and the predicted versus observed concentrations at both the population and individual levels. Furthermore, the final pharmacokinetic model was also evaluated using a visual predictive check.
(VPC), and the uncertainty on each parameter was determined using a non-parametric bootstrap sampling with replacement 1000 times from the original dataset.

2.4 Prediction of Human Clozapine, Norclozapine Brain Concentrations and Expected Receptor Occupancy

After the pharmacokinetic model of clozapine in rat was finalized, the PK model framework was adapted by scaling PK parameters with allometric principles to predict human concentrations in brain. The following exponents were utilized scaling body weight to: clearance 0.75; volume of distribution 1; and first order rate constants 0.25 (Sharma et al., 2009). A 50% of conversion from clozapine to norclozapine in humans was assumed in the model based on prior reports (Centorrino et al., 1994; Raedler et al., 2008; Couchman et al., 2010), and this was implemented in the simulated model assuming $CL_{clo}/F$ is equal to $CL_{clo-p}/F$. The model performance was evaluated comparing model simulated plasma concentrations to published human clozapine concentrations (Ismail et al., 2012) in plasma at steady state following 200, 300 and 400 mg daily doses. The published human clozapine data were reported as total concentrations, and these concentrations were converted to free concentration using 3% unbound fraction (Clozapine Product Insert, 2013) prior to the comparison. After model validation, the simulated human clozapine and norclozapine concentrations were used to calculate the expected human receptor occupancy for multiple receptors. Receptor occupancies were predicted for: dopamine 2(D2); serotonin 2A (5-HT2A); muscarinic-1 (M1); alpha-1 adrenergic (α1); alpha-2 adrenergic (α2); and histamine-1 (H1) using published equilibrium dissociation constants ($K_d$) for clozapine (Seeman et al., 2002; Bumaster et al., 1996; Kroeze et al., 2003) and norclozapine (D2 only) (Lidow et al., 2000).
3. Results

3.1 Rat Population Pharmacokinetics

A two-compartment model with first order absorption best described clozapine pharmacokinetics in rats using a central compartment for plasma concentrations and a peripheral compartment for brain concentrations. Between plasma and brain, an apparent delay in the distribution of clozapine was identified. Several structural models were tested to capture the observed delay. A transit compartment model with two compartments best described flow from plasma to brain, and inter-compartment clearance described the return from brain to plasma (Figure 2.1). Population pharmacokinetic estimates are given in Table 1. Norclozapine exposures in plasma and brain following clozapine administration were adequately described using a similar structure, but with one fewer transit compartment (K\text{tr2}), which was estimated to be approximately 40% of the clozapine value. The volume of distribution of clozapine (V_{clo-p}/F) and norclozapine in brain (V_{met-p}/F) were fixed to the previously estimated values in the final model. Although a significant reduction in the OFV was observed when both parameters were estimated, they were estimated with very poor precision. The elimination of clozapine converted to norclozapine was CL_{clo-met} at 0.055 L/min, which is approximately 10% of total clozapine systemic clearance. The NONMEM control stream with the selected model is also included in the supplementary material (Additional file 1).
Figure 2.1. **Compartmental representation of clozapine and norclozapine pharmacokinetics.**

Two compartments in blue represented the plasma and brain compartment of clozapine.

Two compartments in green represented the plasma and brain compartment of norclozapine.

Clo (Clozapine): $\text{CL}_{\text{clo}}/F$=clearance of clozapine; $V_{\text{clo-p}}/F$=volume of distribution of clozapine in plasma; $K_{a_{\text{clo}}}$=absorption rate of clozapine; $Q_{\text{clo}}/(F\times V_{\text{clo-b}})$=intercompartmental clearance of clozapine; $V_{\text{clo-b}}/F$= volume of distribution of clozapine in brain; $K_{\text{tr1}}$= transit rate constant of clozapine.

Met (Norclozapine): $\text{CL}_{\text{clo}}/F$=clearance of clozapine to norclozapine; $\text{CL}_{\text{met}}/F$=clearance of norclozapine; $V_{\text{met-p}}/F$=volume of distribution of norclozapine in plasma; $K_{a_{\text{met}}}$=absorption rate of norclozapine; $Q_{\text{met}}/(F\times V_{\text{met-b}})$=intercompartmental clearance of norclozapine; $V_{\text{met-b}}/F$= volume of distribution of norclozapine in brain; $K_{\text{tr2}}$= transit rate constant of norclozapine.
<table>
<thead>
<tr>
<th>Parameters</th>
<th>Estimates (RSE%)</th>
<th>BAV (RSE%)</th>
<th>Bootstrap analysis Median [5-95th percentiles]</th>
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<tbody>
<tr>
<td>Clozapine</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>$CL_{clo}/F$ (L/min)</td>
<td>0.5(20.3)</td>
<td>0.463[0.358-0.661]</td>
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<td>$V_{clo-p}/F$ (L)</td>
<td>19.4(40.3)</td>
<td>18.9[5.43-31.59]</td>
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<tr>
<td>$Ka_{clo}$ (1/min)</td>
<td>0.00801(14.7)</td>
<td>0.00815[0.0051-0.0095]</td>
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<td>2.08[0.58-3.43]</td>
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<td>0.20[0.07-0.29]</td>
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<tr>
<td>$Ktr1$ (1/min)</td>
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<td>0.0129[0.011-0.015]</td>
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<tr>
<td>$F_{clo}$</td>
<td>1 FIXED</td>
<td>0.243[0.062-0.35]</td>
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<tr>
<td>Norclozapine</td>
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<td>$CL_{clo-met}/F$ (L/min)</td>
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<td>$CL_{met}/F$ (L/min)</td>
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<td>0.00296[0.0015-0.0047]</td>
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<td>$Ktr2$ (1/min)</td>
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<td>0.00521[0.0045-0.0064]</td>
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<tr>
<td>$F_{met}$</td>
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Residual error (proportional)

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<tr>
<th></th>
<th>Estimates</th>
<th>BAV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parent-plasma</td>
<td>0.109(70)</td>
<td>0.084[0.029-0.19]</td>
</tr>
<tr>
<td>Parent-brain</td>
<td>0.0367(27)</td>
<td>0.0371[0.023-0.055]</td>
</tr>
<tr>
<td>Metabolite-plasma</td>
<td>0.0762(24.1)</td>
<td>0.0714[0.051-0.11]</td>
</tr>
<tr>
<td>Metabolite-brain</td>
<td>0.014(18)</td>
<td>0.0133[0.0095-0.017]</td>
</tr>
</tbody>
</table>

Table 2.1. **Parameter estimates of final population pharmacokinetic model.**

Clo (Clozapine): $CL_{clo}/F$=clearance of clozapine; $V_{clo-p}/F$=volume of distribution of clozapine in plasma; $Ka_{clo}$=absorption rate of clozapine; $Q_{clo}/F$=intercompartmental clearance of clozapine; $V_{clo-b}/F$=volume of distribution of clozapine in brain; $Ktr1$=transit rate constant of clozapine.
Met (Norclozapine): $\text{CL}_{\text{clo-met}} / F=$clearance of clozapine to norclozapine; $\text{CL}_{\text{met}} / F=$clearance of norclozapine; $V_{\text{met-p}} / F=$volume of distribution of norclozapine in plasma; $K_{\text{a,met}} =$absorption rate of norclozapine; $Q_{\text{met}} / F=$intercompartmental clearance of norclozapine; $V_{\text{met-b}} / F=$ volume of distribution of norclozapine in brain; $K_{\text{tr2}} =$ transit rate constant of norclozapine.

BAV=between animal variability

RSE$\%=$ percent relative standard error
Additional file1: NONMEM control stream

...

$SUBS ADVAN13 TOL=6

$MODEL

  COMP(DEPOT) ; ORAL DOSE OF CLOZAPINE
  COMP(CENTRAL) ; PLASMA COMPARTMENT FOR CLOZAPINE
  COMP(PER) ; BRAIN COMPARTMENT FOR CLOZAPINE
  COMP(TRANSIT1) ; TRANSIT COMPARTMENT 1 FOR CLOZAPINE
  COMP(TRANSIT2) ; TRANSIT COMPARTMENT 2 FOR CLOZAPINE
  COMP(MET) ; PLASMA COMPARTMENT FOR NORCLOZAPINE
  COMP(DEPOT2) ; ORAL DOSE OF NORCLOZAPINE
  COMP(PER2) ; BRAIN COMPARTMENT FOR NORCLOZAPINE
  COMP(TRANSIT3) ; TRANSIT COMPARTMENT 1 FOR NORCLOZAPINE

$PK

  CLclo=THETA(1)
  CLclo-met=THETA(2)
  Vclo-p=THETA(3)
  KAclo=THETA(4)
  S2=Vclo-p
  Vclo-b=THETA(5)
  Qclo=THETA(6)*EXP(ETA(1))
  Fclo=THETA(7)*EXP(ETA(2))
  KTR1=THETA(8)*EXP(ETA(3))
  CLmet-p=THETA(9)*EXP(ETA(4))
  KAmet=THETA(10)*EXP(ETA(5))
  Vmet-p=THETA(11)*EXP(ETA(6))
  Vmet-b=THETA(12)
  Qmet=THETA(13)
  KTR2=THETA(14)
  Fmet=THETA(15)

  A._0(1)=0
  A._0(2)=0
  A._0(3)=0
\[ A_0(4) = 0 \]
\[ A_0(5) = 0 \]
\[ A_0(6) = 0 \]
\[ A_0(7) = 0 \]
\[ A_0(8) = 0 \]
\[ A_0(9) = 0 \]

;---------------------------------------------------------------------------------------------------------------------------------

$DES$

;-----------------------------------------------------------------------------PK ODES--------------------------------------------

\[ \text{DADT}(1) = -K\text{Aclo} \times A(1) \]
\[ \text{DADT}(2) = K\text{Aclo} \times A(1) - (\text{CLclo-met/Vclo-p}) \times A(2) - (\text{CLclo/Vclo-p}) \times A(2) - (Q\text{clo/Vclo-b}) \times A(3) - K\text{TR1} \times A(2) \]
\[ \text{DADT}(3) = K\text{TR1} \times A(5) - (Q\text{clo/Vclo-b}) \times A(3) \]
\[ \text{DADT}(4) = K\text{TR1} \times A(2) - K\text{TR1} \times A(4) \]
\[ \text{DADT}(5) = K\text{TR1} \times A(4) - K\text{TR1} \times A(5) \]
\[ \text{DADT}(6) = (\text{CLclo-met/Vclo-p}) \times A(2) - (\text{CLmet/Vmet-p}) \times A(6) + K\text{Amet} \times A(7) + (Q\text{met/Vmet-b}) \times A(8) - K\text{TR2} \times A(6) \]
\[ \text{DADT}(7) = -K\text{Amet} \times A(7) \]
\[ \text{DADT}(8) = K\text{TR2} \times A(9) - (Q\text{met/Vmet-b}) \times A(8) \]
\[ \text{DADT}(9) = K\text{TR2} \times A(6) - K\text{TR2} \times A(9) \]

...
Figures 2.2 and 2.3 show the goodness of fit plots of the final model for the parent drug and metabolite in plasma and brain, respectively. Population and individual predictions as well as the conditional weighted residuals distribution are shown in these figures. In the case of norclozapine, imprecision in population predicted plasma concentrations was evident and is attributed to the inter-animal variability observed in the context of the limited number of animals available to support these predictions. The majority of the fixed effects were estimated with less than 40% relative standard error (Table 1). BAV was estimated for several of the structural parameters and ranged from 5% (K_{tr1}) to 75% (Cl_{met}). Residual variability for clozapine in plasma and brain were 10.9% and 3.7%, respectively, and residual variability of norclozapine in plasma and brain were 7.6% and 1.4%, respectively. VPC results are shown in Figures 2.4 and 2.5, the observed medians (dashed black lines) concentrations were adequately captured by the corresponding simulation based 90% predicted intervals of median concentrations for clozapine and norclozapine (shaded areas). Median, 5\textsuperscript{th} and 95\textsuperscript{th} percentiles of the parameters derived from the bootstrap analysis of 1000 replicates are shown in Table 2.1.
Figure 2.2. Model diagnostic plots for clozapine.

A and E: population prediction vs. observation plots for clozapine in plasma and brain ECF, respectively, solid line is line of identity. B and F: individual prediction vs. observation plots for clozapine in plasma and brain ECF, respectively. C and G: conditional weighted residuals vs. time for clozapine in plasma and brain ECF, respectively. D and H: conditional weighted residuals vs. population prediction for clozapine in plasma and brain ECF, respectively.
Figure 2.3. Model diagnostic plots for norclozapine.

A and E: population prediction vs. observation plots for norclozapine in plasma and brain ECF, respectively, dashed line is the line of identity. B and F: individual prediction vs. observation plots for norclozapine in plasma and brain ECF, respectively. C and G: conditional weighted residuals vs. time for norclozapine in plasma and brain ECF, respectively. D and H: conditional weighted residuals vs. population prediction for norclozapine in plasma and brain ECF, respectively.
Figure 2.4. **Visual predictive checks of clozapine in plasma and brain and norclozapine in plasma.**

Visual predictive check of clozapine concentrations in plasma (A) and brain (B), and norclozapine concentrations in plasma (C) following a 10 mg/kg subcutaneous dose of clozapine. Norclozapine concentrations in brain following a 10 mg/kg subcutaneous dose of clozapine were not measureable.

Dashed line is the median of observed concentrations, the shape represents the 90% predicted interval of the median, and the dots represent the observed concentrations.
Figure 2.5. **Visual predictive checks of norclozapine in plasma and brain.**

Visual predictive check of norclozapine concentrations in plasma (A) and brain (B) following a 10 mg/kg subcutaneous dose of norclozapine.

Dashed line is the median of observed concentrations, the shape represents the 90% predicted interval of the median, and the dots represent the observed concentrations.
3.2 Human PK Simulation

Simulated unbound clozapine concentrations in plasma from 12 to 24 hours after administration were compared with published human data and are shown in Figures 2.6A, 2.6B and 2.6C for 3 doses (200, 300 and 400mg/day). Predicted occupancy of D2, 5-HT2A, M1, α1, α2 and H1 receptors were calculated using simulated unbound clozapine and norclozapine (D2 only) brain concentrations for the three dose levels. For clozapine, the predicted median percentage of receptor occupancy of D2 ranged from 6-42%, 9-52% and 11-59% for the 200, 300 and 400mg daily doses, respectively, across the inter-dose time interval. The median percentage of 5-HT2A receptor occupancy decreased from 93% to 52%, 95% to 62% and 96% to 69% from 6 to 24 hours after 200, 300 and 400mg daily doses, respectively. For M1, α1 and H1 receptors, occupancies ranged from 74 to 99% across the dosage interval. In addition, the median percentage occupancy of α2 receptors was predicted to be in the range of 3-40% across the dosage interval. For norclozapine, the predicted median percentage of receptor occupancy of D2 ranged from 1.1-17.3% across the dosage interval. Receptor occupancy results across the dosage interval are summarized in Figure 2.7.
Figure 2.6. Simulated clozapine human unbound concentrations vs. published human concentrations at steady state in brain with 200, 300 and 400 mg OID from 12 to 24 hours. The shape represents the 90% predicted interval of the median and the dots represent the observed data.
Figure 2.7. The predicted median percentage of receptor occupancy of D2, 5-HT2A, M1, α1, α2 and H1 for clozapine and D2 for norclozapine.

The predicted median percentage of receptor occupancy of D2, 5-HT2A, M1, α1, α2 and H1 for clozapine were shown in A to F, respectively, and predicted median percentage of receptor occupancy of D2 for norclozapine was shown in G between 6 to 24 hour after dose. The solid line, dash line and dot line represent the predicted median percentage of receptor occupancy following 200, 300 and 400 mg daily doses, respectively.
4. Discussion

The model presented represents a unique PK model developed from directly measured concentrations of clozapine and norclozapine in rat plasma and brain ECF. A multiple transit compartment model was used to account for a delay in the transport of clozapine and norclozapine from plasma to brain across the rat blood brain barrier. Some evidence suggests that Pgp may be involved in the process of clozapine transport (Cremers et al., 2012; Doran et al., 2005) across the blood brain barrier. The need to incorporate transit compartments in the present model is consistent with a Pgp role in clozapine transport across this barrier. Using an animal model, drug exposure can be measured by microdialysis at the target site. Based on a previously published non-compartmental analysis, the ratio of AUC between parent and metabolite in the rat indicated that only about 10% of parent drug was eliminated through metabolism (Olsen et al., 2008). This is consistent with the ratio of norclozapine to clozapine clearance ($Cl_{\text{clo-met}}$ is 9.91% of $Cl_{\text{clo}}$). As the results revealed, even with relatively rich sampling profiles, the uncertainty of some parameters, in particular of between-animal variability, was large likely because of the small number of animals in this study.

Simulated human plasma concentrations were based on previously published human plasma concentration data (Ismail et al., 2012). The unbound plasma concentrations at steady state after a range of doses overlapped with published data corrected for the unbound fraction of clozapine (3%) (Clozapine Product Insert, 2013). Subsequently, plasma exposures were linked to the plasma–brain structural PK parameters, using allometric scaling, that described clozapine and norclozapine transport between plasma and brain in the rat to ultimately predict human brain ECF exposure. This population
pharmacokinetic approach, based on a transit compartmental approach as opposed to explicit assumption of a Pgp role and its associated interspecies scaling, enabled translational representation of the system across species to predict human brain ECF concentrations. As an atypical antipsychotic drug, clozapine targets D2 receptors as well as acts as an agonist or antagonist at several other receptors found in the CNS. In order to get a more complete profile of PK-PD linkage, percentage receptor occupancy of each receptor was calculated from 6 to 24 hour after three dosage levels. Our results show that the median percentage D2 receptor occupancy was in a range of 42% to 59% 6 hours after administration of a daily dose of 200-400 mg. This range is congruent with the 33% to 67% range reported by Nordström et al. (Nordström et al., 1995), and agrees with the widely recognized understanding of low D2 receptor occupancy of therapeutic doses of clozapine relative to those obtained with therapeutic doses of other antipsychotics (first and second generation). In addition to D2, percent 5-HT2a receptor occupancy also overlapped with the results of Nordström et al.

The proposed PK model thus demonstrated the ability to extrapolate human systemic exposure to predict clozapine brain concentrations and associated receptor occupancy profiles in humans at clinically relevant doses. In addition, the model simultaneously captured parent and metabolite in the system, which is relevant since norclozapine also has activity at multiple receptors (Bishara et al., 2008). However, the model can be improved in the precision of the PK parameter estimates by increasing the sample size. With this limitation taken into consideration, the model framework reported shows promise in predicting clozapine receptor occupancy at multiple receptors in human CNS, which can then be probed as a correlate to response and/or toxicity.
CHAPTER III: Population Pharmacokinetic Modeling of Sertraline Treatment in Alzheimer’s disease Patients: The DIADS-2 Study (Published in The Journal of Clinical Pharmacology) (This material is reproduced with permission of John Wiley & Sons, Inc.)

1. Introduction

Alzheimer’s disease (AD) is a neurodegenerative disease associated with a number of neuropsychiatric symptoms (NPS). One commonly found NPS is depression, affecting as many as 60% of AD patients (Steinberg et al., 2008). The antidepressant sertraline, a selective serotonin reuptake inhibitor (SSRI), has been used for the treatment of depression in AD patients (Martin et al., 2006). It is the second most potent inhibitor of serotonin reuptake (Hiemake et al., 2000). In a study of 117 randomized controlled trials from 1991 to 2007, sertraline was proposed as the best first line treatment for moderate to severe depression in adults based on an overall evaluation of benefits, acceptability and other factors (Cipriani et al., 2009). Sertraline is orally administered with high plasma protein binding affinity (Owen et al., 1997). The average elimination half-life of sertraline is approximately 26 hours and the peak plasma concentration (C_{max}) is reached at 6-8 hours (Warrington et al., 1991). Sertraline is mainly eliminated by hepatic metabolism to its major metabolite, N-desmethylsertraline, by multiple cytochrome p450 enzymes including CYP2B6, CYP2D6, CYP2C9, CYP2C19 and CYP3A4 (Kobayashi et al., 1999). This metabolite has 5 -10% of sertraline’s serotonin reuptake inhibitor potency; thus its clinical effect on sertraline response is negligible (Sprouse et al., 1996).

The pharmacokinetic profile of sertraline has been broadly explored in previous clinical studies where patient ages spanned broad ranges (Muijsers et al., 2002; Schneider et al., 2003; Axelson et al., 2002). In a pharmacokinetic study of 16 elderly (≥65 years of
In the elderly, sertraline’s effectiveness is comparable to the SSRI fluoxetine as well as the tricyclic antidepressants (TCAs) nortriptyline, amitriptyline and imipramine. It has lower rates of adverse side effects than the TCAs (Muijsers et al., 2002). Although many studies have examined the sertraline pharmacokinetic profile in elderly subjects, non-compartmental methods were employed that have limitations in assessing sources of inter-individual variability in sertraline concentration. In fact, this is the first population pharmacokinetic (PPK) study focusing on AD patients with depression. While this analysis was based on data from a null clinical study, it provided an opportunity to capture the pharmacokinetic characteristics in elderly individuals of sertraline. In these analyses, we aim to gain insights relating to inter-individual variability in the pharmacokinetics of sertraline in AD patients. The objective is to identify covariates that contribute to variability in sertraline concentration by performing a PPK analysis of sertraline in elderly patients with AD and generating PPK parameters for this population.

2. Materials and Methods

2.1 Participants and Study design

The design of the multicenter Depression in Alzheimer’s Study-2 (DIADS-2) has been described in detail elsewhere (Martin et al., 2006; Rosenberg et al., 2010; Weintraub
et al., 2010; Drye et al., 2011). Briefly, DIADS-2 enrolled 131 AD patients with mild-to-moderate AD. Patients were randomized in a 12-week, double-blind, placebo-controlled (n=64) antidepressant trial of sertraline (n=67; range: 25-125 mg per day). An initial treatment regimen of sertraline 50 mg QD or identical-appearing placebo was prescribed. The dosage of sertraline in the active treatment arm was increased to 100 mg QD after one week. The daily dose was adjusted depending on the response and tolerability of the treatment in the first four weeks post randomization. Single concentration samples of sertraline were collected in individual patients at weeks 4 and 12. The time of last dose and exact time of collection were available for each of these samples.

The study was approved by Institutional Review Boards at all five study sites and the coordinating center: Johns Hopkins School of Medicine, Baltimore, MD; University of Pennsylvania School of Medicine, Philadelphia, PA; Medical University of South Carolina, Charleston, NC; University of Rochester School of Medicine, Rochester, NY; University of Southern California Keck School of Medicine, Los Angeles, CA; Johns Hopkins Bloomberg School of Public Health. In addition, the PPK analysis was approved by the institutional Review Board of Indiana University School of Medicine, Indianapolis, IN.

2.2 Analytical Procedure to Measure Sertraline Concentration

Plasma sertraline concentration was determined using high-performance liquid chromatography (HPLC). The extraction of plasma was done in the mobile phase at a 60:40 ratio of 0.025 M potassium phosphate buffer, pH 2.5, and acetonitrile with a flow rate of 1 mL/min. The separation of plasma occurred in a Knauer nucleosil, C18, 100 angstrom, 150 mm x 4.6 mm column with a Supelco pelliguard LC-18, 2 cm X 4.6 mm
pre-column cartridge. Ninety microliters of n-octylamine was added as a modifier to the mobile phase and degassed. The sertraline and D-sertraline assays were linearized by an internal standard, 200 ng/mL of clomipramine, in the range of 10 to 600 ng/mL. The intra-assay and inter-assay variability in the coefficient of variation (CV) ranged from 6.6% to 9.4% and ranged from 2.8% to 6.3%, respectively. With 1 mL of plasma, recovery for sertraline and metabolite ranged from 87% to 93% with a lower limit of quantification (LLOQ) of 10ng/mL. The analysis was carried out using a Turbo Chrom data system.

2.3 Population Pharmacokinetic Model Development

The population pharmacokinetics of sertraline were analyzed using nonlinear mixed effect modeling software, NONMEM, Version VII (GloboMax LLC, Ellicott City, MD, USA) using Wings for NONMEM, Version 7 (Holford, 2012). The initial model development focused on a base model structure on PK parameter assessment. One and two compartment models with first order absorption and elimination were evaluated using subroutine ADVAN2 TRANS2 and ADVAN4 TRANS4, respectively. A likelihood based approach (Method 3) was used to handle measurements below the quantitation limit (BQL) at 10 ng/mL (Ahn et al., 2008).

PPK analyses used the first-order conditional estimation (FOCE) LAPLACIAN method. Inter-individual variability (IIV) for PK parameters was assumed to be log-normally distributed and evaluated using an exponential model $P_i = P_{TV} \times e^{\eta_p}$ where $P_i$ is the parameter estimate for the ith individual, and $P_{TV}$ is the typical value for the parameter at the population level. The variability between ith individual and population parameter values was described by $\eta_p$ which was identically distributed with a mean of 0 and a
variance of $\omega_\eta^2$ (Feng et al., 2006). In addition to the IIV, intraindividual variability, system noise, experimental error and/or model misspecifications was described by a residual error model. The residual error models evaluated were: additive ($y_{ij} = \hat{y}_{ij} + \varepsilon_{ij}$); proportional ($y_{ij} = \hat{y}_{ij}(1 + \varepsilon_{ij})$); and combined ($y_{ij} = \hat{y}_{ij}(1 + \varepsilon_{ij}) + \varepsilon_{ij}'$); where $y_{ij}$ and $\hat{y}_{ij}$ represents the jth observed sertraline concentration, and its corresponding model predicted concentration with the difference described by $\varepsilon_{ij}$ or $\varepsilon_{ij}'$. $\varepsilon_{ij}$ was assumed to be normally distributed with a mean of 0 and a variance of $\sigma^2$. The absorption rate constant (Ka) was fixed to 0.5 based on the literature values of $t_{\text{max}}$ and an elimination constant (Ronfeld et al., 1997). This was done because the estimation of Ka in this dataset resulted in unstable model runs.

To evaluate the inter-individual variability estimated by the nonlinear mixed effects modeling approach, patients’ demographic characteristic (weight, height, age, sex, race and study site) were evaluated to see if these explained this variability. These factors were assessed independently in a step-wise forward addition approach. Covariates such as weight, height, age, sex and race were included to examine potential physiologic differences that could contribute to difference in drug elimination rate or distribution volume across this population. Given the possibility of differences in adherence to the protocol by either subjects or the study site in sample collections, etc, clinical site was also tested as a covariate.

For continuous covariates, the effects of the covariates on PK parameter estimates were tested in the following model structures:

$$(1) \ P_{\text{TV}} = \theta_1 + \theta_2 * \text{Cov}$$

$$(2) \ P_{\text{TV}} = \theta_1 + \theta_2 * (\text{Cov} - \text{Med}_{\text{cov}})$$
(3) $P_{TV} = \theta_1 \times (Cov / Med_{cov})^{0.2}$

(4) $P_{TV} = \theta_1 \times \exp[\theta_2 \times (Cov / Med_{cov})]$

where $P_{TV}$ is the typical population estimate of a particular PK parameter, and $\theta_1$ and $\theta_2$ are fixed effect estimates for a corresponding covariate, Cov normalized by the median value of the covariate, Med$_{cov}$. Missing data were found in both weight and height covariates. A naïve substitution approach (Yang et al) was carried out to simply replace the missing value with the median value based on sex.

(5) IF (WT=0 and SEX=1) THEN WT= Med$_{WT,Sex=Male}$

IF (WT=0 and SEX=2) THEN WT= Med$_{WT,Sex=Female}$

A common allometric function of scaling the PK parameters to the 0.75 power of body weight was also tested after the missing weight values were replaced.

(6) $P_{TV} = \theta_1 \times (WT / Med_{WT})^{0.75}$

Categorical variables such as sex, race and site were tested in the following model structure. Each category was evaluated in a separate fashion.

(7) IF (Cov.EQ.1) THEN $P_{TV} = \theta_1$

Else $P_{TV} = \theta_2$

For example, each ethnic group was divided into a category, African American=1, White=4 and Hispanic/Latinos=5. Each ethnic group was examined separately from other races. In addition, groupings of racial categories were affected when the individual race effects were not uniquely identifiable. In this case, $\theta_1$ and $\theta_2$ are the population PK parameter estimates for groupings of race that were uniquely identifiable, in this case African American and all others.
Model evaluation was based on a likelihood ratio test using the objective function value (OFV) from NONMEM. The change in the OFV returned by NONMEM is approximately equal to -2 x log likelihood. The difference in -2 x log likelihood between two models that are nested follows a $\chi^2$ distribution. Covariates were added to the model in a step-wise addition fashion and remained in the final model if the OFV decreased by greater than 3.84 (p-value $\leq 0.05$, df=1). The final model was further examined using goodness-of-fit plots generating using R (Version 2.13) based on the conditional weighted residuals distribution and the predicted versus observed sertraline concentrations at both the population and individual levels.

3. Results

3.1 Patient Characteristics

A total of 131 participants entered the trial with 67 randomized to sertraline and 64 to placebo. Only the concentration measurements taken from patients in the active treatment arm (n=67) were utilized for this analysis. An average of 1.7 sertraline concentration measurements per individual was available, and 5 of the measurements were found below the LLOQ. Seventeen individuals were removed from the analysis. Specifically, 16 individuals lacked sertraline concentration information at both weeks 4 and 12, and one individual had missing dosage information for both occasions. In the remaining 50 individuals, other single observations were removed as follows: 14 individuals only had a single sertraline concentration measurement from one of the two visits (non-measured visit removed). Of these 14 observations, 11 were missing a concentration measurement, 1 was missing a dosage time associated with a concentration sample, and 2 were missing dosage amount information associated with that
concentration sample. The PPK analysis was conducted using the remaining 85 PK observations from 50 individuals.

As shown in Table 3.1, the median age of the patients was 75 (range: 53-89). There were 20 males and 30 females in the analysis broken down as follows: 18% African American; 72% White; and 10% Hispanic. This analysis included 4 patients with missing weight information and 9 with missing height information. The median values of weight and height without considering the missing values were 147 lb (range: 107-245) and 64 in (range: 57-72), respectively.
Table 3.1. Alzheimer’s disease patient demographics

<table>
<thead>
<tr>
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<th>N (%)</th>
<th>Mean (SD)</th>
<th>Median(range)</th>
</tr>
</thead>
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<td>Sample size</td>
<td>50</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of observations</td>
<td>85</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gender</td>
<td></td>
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<tr>
<td>Male</td>
<td>20 (40)</td>
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<td></td>
</tr>
<tr>
<td>Female</td>
<td>30(60)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Race</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Black/African American</td>
<td>9 (18)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>White</td>
<td>36 (72)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hispanic/Latino</td>
<td>5 (10)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clinical sites</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>15 (30)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B</td>
<td>11 (22)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>6 (12)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>D</td>
<td>8(16)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>E</td>
<td>10(20)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline Age (years)</td>
<td></td>
<td>75 (7.76)</td>
<td>76 (53-89)</td>
</tr>
<tr>
<td>Baseline Weight (Ib)</td>
<td></td>
<td>159 (37.08)</td>
<td>147 (107-245)</td>
</tr>
<tr>
<td>Without 4 missing values</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline Height (In)</td>
<td></td>
<td>64 (4.45)</td>
<td>64 (57-72)</td>
</tr>
<tr>
<td>Without 9 missing values</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sertraline dose administered (mg)</td>
<td>92.47 (18.62)</td>
<td>100 (25-100)</td>
<td></td>
</tr>
<tr>
<td>Sertraline concentrations (ng mL⁻¹)</td>
<td>62.94 (47.62)</td>
<td>49 (9-229)</td>
<td></td>
</tr>
<tr>
<td>Without 5 BQL values</td>
<td></td>
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</tr>
</tbody>
</table>
3.2 Population Pharmacokinetic Modeling

A one-compartment model with first order absorption and elimination and an additive residual error model best described the sertraline data. The population parameter estimates of CL/F and Vd/F in the base model were 83.1 L/h and 6,620 L, respectively. Inter-individual variability (IIV) was estimated only for CL/F because a significant correlation was found between CL/F and Vd/F. Patients at site C has CL/F approximately 49% lower than that seen in patients other at other clinical sites ($\chi^2 = 5.576$ df=1, p<0.05). The final covariate model was implemented using the categorical covariate model structure described in the methods section (equation 7). No other significant covariate relationships were found for CL/F or Vd/F. The population PK parameter estimates and goodness-of-fit plots for the final model are listed in Table 3.2 and Figure 3.1, respectively.
Table 3.2. **Population pharmacokinetic parameter estimates of sertraline in the final model**

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Population estimate (% SE)</th>
<th>Inter-individual variability (% SE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CL/F_site=C, L/h</td>
<td>43.8(34)</td>
<td>59.33% (31.3)</td>
</tr>
<tr>
<td>CL/F_site=others, L/h</td>
<td>89.1(12.2)</td>
<td>59.33% (31.3)</td>
</tr>
<tr>
<td>Vd/F, L</td>
<td>6470(70.5)</td>
<td></td>
</tr>
<tr>
<td>Ka, 1/h (fixed)</td>
<td>0.5</td>
<td></td>
</tr>
<tr>
<td>Residual variability</td>
<td>19.6 ng/mL (11.6)</td>
<td></td>
</tr>
</tbody>
</table>

CL/F\_site=C; clearance adjusted for bioavailability from site C; CL/F\_site=others; clearance adjusted for bioavailability from other clinical sites; Vd/F, Volume of distribution adjusted for bioavailability; Ka, rate of absorption; SE, standard error
Figure 3.1. Goodness-of-fit plots for the final sertraline model.

Plot of population and individual predicted versus observed sertraline concentrations and plot of conditional weighted residuals versus population predicted concentrations and time (hours).
4. Discussion

Nineteen percent of the variance in pharmacokinetic distribution of sertraline in these depressed AD patients was accounted for using a 1-compartment model. The population mean CL/F and Vd/F were 83.1 L/h and 6,620 L, respectively. As most patients in DIADS-2 were elderly (mean age of 75 years; range: 53-89), the mean elimination rate constant of 0.013 1/h is consistent with other literature reported values (Ronfeld et al., 1997). Compared to younger patients, the elderly have a longer sertraline half-life, but we found no difference between elderly healthy volunteers in the literature and these AD patients.

In our analysis, CL/F estimates for the four patients with sertraline observations below the quantification limits were in the top 24% of all CL/F estimates. Indeed, one patient with both observations below the quantification limit had a CL/F value approximately three times higher than the typical population value. The cause of these high CL/F values is not clear. Many possible factors might be considered, such as fasting, poor adherence, or genetic variance.

We also examined the effect of sex, race, age, weight, height and site on the variability of the PK parameter estimates. Previous publications have suggested that the average half-life is 1.5 times longer in women than in men (Ronfeld et al., 1997). We expected to detect an effect of gender on sertraline PK parameters; however, inclusion of sex as a covariate in our model yielded a statistically non-significant association. Possible differences between males and females were likely undetectable because of the small study population. In addition, a commonly reported correlation (Mahmood et al., 2007) between plasma clearance and body weight was not detected in this analysis.
Unexpectedly, the only covariate examined that affected PK variability was related to a single clinical site. Patients at site C had much lower CL/F than other sites. This might be explained by the 3 high sertraline concentrations found in 2 patients out of a total of 9 observations from site C. Nineteen percent of the total variance in inter-individual on CL/F was explained by incorporating the site as a covariate in this analysis. The model CL/F estimates we present that exclude patients at the site C are likely more generalizable to the typical AD population. In conclusion, clinical site was a significant covariate contributing to the CL/F change. The clinical implications for subjects at the outlying site would be important if this represented a true bias in subject selection. In this case diminished clearance could lead to a greater risk of adverse effects such as dizziness, extrapyramidal effects and hyponatremia. Nonetheless, the subjects at this site were found not to differ in demographic nor clinical attributes from the other four sites. This, as well as the failure to detect differing incidences of side effects, leads to the more likely explanation, that there may have been variations in study procedures such as sampling.

Site variability in trial procedures is critical to the validity of multi-site trials, and population pharmacokinetics may prove helpful in this assessment. If drug administration and sample collection procedures were found not to be atypical than this may suggest meaningful subject differences in the patient sample at the outlying site e.g., drug metabolism or body size as we have found previously (Jin et al., 2010). Therefore, the dosage regimen and administration should be closely monitored in a multicenter study in order to avoid unnecessary exposures or incomplete treatments.
CHAPTER IV: Data quality constrains utility of computational modeling of tumor burden in non-small cell lung cancer clinical trials

1. Introduction

Cancer drug development would be more effective if new therapeutics could be evaluated with readily available technologies, in fewer patients, observed on treatment for shorter periods of time. One set of strategies to achieve these goals has been computational modeling of the longitudinal growth of non-small cell lung cancer (NSCLC) in populations of patients and in silico simulation of clinical trials. (Claret et al., 2012; Houk et al., 2009; Wang et al., 2009; Fridlyand et al., 2011; Moertel et al., 1976). The ultimate goal of these efforts is to improve the efficiency of cancer drug clinical development (Barrett et al., 2007; Bruno et al., 2009; Bruno et al., 2010).

In NSCLC, clinical trial simulations have employed a “two-step” joint model (Ibrahim et al., 2010) of overall survival in metastatic disease that was based on a longitudinal tumor growth model. These models were developed with data from nearly 3,400 patients submitted to the FDA. From 4 randomized phase III clinical trials for regulatory approval of bevacizumab, docetaxel, erlotinib, and pemetrexed (Wang et al., 2009) the longitudinal growth model was derived from the sum of the longest dimension measurements of tumors by computed tomography (CT) imaging as recorded in study case report forms. Interpolations of the change in tumor size from baseline to 8 weeks of treatment (the tumor size ratio or TS) proved an important predictor of overall survival. Modeling and simulation with these data could be an efficient means to support decision making at the Phase 2 to Phase 3 transition in drug development (Claret et al., 2012).
Another potential benefit of quantitative analysis of NSCLC tumor burden would be to redesign phase II trials to randomize fewer patients and have shorter observation periods than required for determining progression-free survival (Adjei et al., 2009; Dhani et al., 2009; Maitland et al., 2011; Maitland et al., 2011; Mandrekar et al., 2010; Stein et al., 2011; Yap et al., 2010). A simple strategy in NSCLC would be to measure the median change in tumor size at 8 weeks for randomly-assigned treatments among the study arms (Bruno et al., 2009; Lavin et al., 1981; Karrison et al., 2007). A more complex strategy that might ultimately require fewer resources would be to collect all tumor measurements from all patients over time and have the cumulative data from patients enrolled early in the trial continuously inform a calculated parameter of drug effect. These strategies have had limited testing and require validation. For example, in studies of colorectal cancer therapy and survival outcomes some have found advantages to continuous tumor measurement metrics while others have not (An et al., 2011; Kaiser et al., 2013; Claret et al., 2013).

The purpose of this study was to assess and refine the published FDA longitudinal tumor size model for NSCLC using archived tumor measurement data so that modeling and simulation might lead to smaller, quicker early phase trials for testing new treatments for NSCLC. We obtained archived case report forms from three randomized clinical trials by the Cancer and Leukemia Group B (CALGB) sponsored by the U.S. National Cancer Institute (NCI). We intended to evaluate the power of smaller clinical trials with novel endpoints to detect evidence of anti-cancer drug treatment effects. Instead, we found elements of 3 data sets from multi-center clinical trials that could bias comparisons between continuous measurement and categorical strategies for improving treatment
evaluations. These findings are likely to be common to historical and current solid tumor trial data sets. Here we identify variance in measurement and recording of CT imaging assessments of tumor burden as a modifiable factor that constrains the successful development and validation of novel tumor growth assessment metrics.

2. Materials and Methods

2.1 Patients

Archived case report forms were available from 857 patients enrolled in 3 NCI-supported studies conducted by CALGB listed in Table 4.1. These were front-line trials in metastatic NSCLC: CALGB 9730 (Lilenbaum et al., 2005) was a phase III randomized trial that compared single-agent paclitaxel with combination carboplatin/paclitaxel, CALGB 30203 (Edelman et al., 2008) was a randomized phase II trial that evaluated eicosanoid modulation and examined cyclooxygenase-2 expression as a positive predictive factor for the inclusion of celecoxib in the standard first-line cytotoxic therapy regimens, and CALGB 30303 (Miller et al., 2008) was a phase II randomized study of dose-dense docetaxel and cisplatin administered every two weeks with pegfilgrastim and darbopoetin-alfa with or without the chemoprotectant BNP7787. The inclusion and exclusion criteria of the trials were previously published (Edelman et al., 2008; Lilenbaum et al., 2005; Miller et al., 2008).
Table 4.1. **Three U.S. National Cancer Institute (NCI)-sponsored studies conducted by the Cancer and Leukemia Group B CALGB**

<table>
<thead>
<tr>
<th>CALGB study</th>
<th>Treatment</th>
<th># of patients enrolled</th>
<th># of patients treated &amp; eligible</th>
<th>Dates of accrual</th>
</tr>
</thead>
<tbody>
<tr>
<td>30203</td>
<td>carboplatin/gemcitabine + zileuton/celecoxib/both</td>
<td>140</td>
<td>134</td>
<td>12/2003 – 9/2004</td>
</tr>
</tbody>
</table>
2.2 Original Clinical Trial Data Collection

Data relevant to original reporting of the clinical trial results were captured on case report forms and entered into the CALGB digital databases for each of the clinical trials. The coded, patient-level data were stored at the Core Statistical Facility for CALGB (Durham, NC, USA). Treatment response assessments were conducted according to the study protocols. The CALGB 9730 trial incorporated standard World Health Organization response criteria (Miller et al., 1981) based on imaging studies conducted every 2 cycles (6 weeks). Bi-dimensional measurements of radiographically identified lesions were typically hand-written into the same data-field on a standard spreadsheet at each evaluation time-point. Treatment was discontinued: when it became intolerable; when new lesions were identified; when the product of two perpendicular diameters of any measured lesion increased by at least 25%; or when 6 cycles of therapy were completed. Patients were subsequently followed every 3 months for 2 years. For CALGB studies 30203 and 30303, the Response Evaluation Criteria in Solid Tumors was employed, and categorical responses were based on the sum of the longest unidimensional measurements of criteria-defined “target lesions” (Therasse et al., 2000). Thoracic CT imaging evaluations were conducted in all patients pre-treatment, and at 6 and 12 weeks after treatment. Patients were removed from the studies for unacceptable toxicity or progression of disease. Patients who completed all study therapy were followed at minimum every 12 weeks thereafter.

For studies 30203 and 30303, tumor measurements were recorded on the “CALGB Solid Tumor Evaluation Form” – C-660. The form designates separate rows for each target lesion and columns for each evaluation time-point. The longest unidimensional
measurement was entered for each designated target lesion at baseline in a separate row. Non-target lesions were identified as present in a separate section of the form. During the studies the original case report forms were transmitted by the sites to the CALGB Statistical Facility where the forms were reviewed and verified by the data management team and the RECIST categorical evaluations were entered into digital databases. The sum of the longest dimensions of target lesions and the target lesion measurements were not captured in the study database. After publication of study results, the case report forms and associated source document CT reports were stored off-site in an archive facility.

2.3 Tumor Measurement Collection

To obtain the original target lesion measurements, the archived paper case report forms were obtained from storage, scanned, and saved as portable document format (pdf) files. Tumor measurements from the pdf files for CALGB 30203 and 30303 were manually extracted by a research assistant and entered into a tracking file and into the study databases simultaneously. The transcriptions were independently reviewed by one of the study authors (SK and CL) and inconsistencies were manually corrected. Additionally, individual patient tumor growth plots were inspected for atypical growth and response patterns. All aberrant plots were cross-verified with the original case report form pdf and any additional data entry errors captured by this review were corrected before modeling analyses were performed. This process (Figure 4.1) resulted in 103 patients with evaluable data from CALGB 30203 and 124 from CALGB 30303. A sample of case report forms from CALGB 9730 (Figure 4.2), consistently revealed insufficient documentation of quantitative measurements of tumors to be useful for this
modeling exercise, and therefore all patients from this trial were excluded from our planned evaluation.

Figure 4.1. Selection of patients contributing images from CALGB 30203 and 30303.
Figure 4.2. An example of a case report form from CALGB 9730 trial.

<table>
<thead>
<tr>
<th>Anterior RUL nodule</th>
<th>1.1</th>
<th>nc</th>
<th>nc</th>
<th>nc</th>
<th>nc</th>
<th>nla</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anterior med RML nodule</td>
<td>1.5×1.5</td>
<td>nc</td>
<td>nc</td>
<td>nc</td>
<td>nc</td>
<td>nla</td>
</tr>
<tr>
<td>LLL tissue mass</td>
<td>present</td>
<td>nc</td>
<td>nc</td>
<td>nc</td>
<td>nc</td>
<td>nla</td>
</tr>
<tr>
<td>Tiny pulmonary nodules</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>present</td>
<td></td>
</tr>
<tr>
<td>Uptake proximal R 10th rib</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>present</td>
</tr>
</tbody>
</table>
2.4 Tumor Size Modeling

Longitudinal tumor size trajectories (sum of longest tumor diameter) were analyzed with nonlinear mixed effect modeling software, NONMEM, Version VII (GloboMax LLC, Ellicott City, MD, USA) using Wings for NONMEM, Version 7 (Holford, 2012) and the published model structure (Wang et al., 2009). This model employed a combination of a linear growth function and an exponential shrinkage function to describe the tumor change respect to baseline size (Eq.1).

\[ T_{Si}(t) = BASE_i e^{-SR_i t} + PR_i t \]

Where \( T_{Si}(t) \) is the tumor size at time \( t \) for the \( i^{th} \) individual, \( BASE_i \) is the baseline tumor size, \( SR_i(t) \) is the exponent tumor shrinkage rate constant, and \( PR_i(t) \) is the linear tumor growth rate constant. The exponential shrinkage function constrains the tumor size to be greater than zero (to avoid negative tumor sizes being predicted) and reflects the drug effect on tumor (whether or not there is actual tumor shrinkage). Tumor size changes were modeled using the first-order conditional estimation (FOCE) and stochastic approximation expectation maximization (SAEM) method with interaction. Between subject variability was assumed to be log-normally distributed and evaluated on baseline tumor size, tumor shrinkage rate and tumor progression rate using an exponential model

\[ P_i = P_{TV} \times e^{\eta p} \]

where \( P_i \) is the parameter estimate for the \( i^{th} \) individual and \( P_{TV} \) is the typical value for the parameter at the population level. The variability between \( i^{th} \) individual and population parameter values was described by \( \eta_p \) which was identically distributed with a mean of 0 and a variance of \( \omega^2 \) (Feng et al., 2006). Residual variability was also estimated using a proportional residual error model (\( y_{ij} = \hat{y}_{ij} (1 + \epsilon) \)) where \( y_{ij} \) and \( \hat{y}_{ij} \) represents the \( j^{th} \) observed tumor trajectory, and its corresponding
model predicted tumor size. The difference between observed and predicted values was described by $e_{ij}$ which was assumed to be normally distributed with a mean of 0 and a variance of $\sigma^2$.

A likelihood ratio test was applied based on the change in the objective function value (OFV) in NONMEM to evaluate the random effect on parameters. This change in OFV is approximately equal to $-2 \times \log \text{likelihood}$. The difference in $-2 \times \log \text{likelihood}$ between two models that are nested follows a $\chi^2$ distribution. An alpha level for significance of 0.05 was set a priori (3.84 OFV for 1 degree of freedom). The final model was further examined using goodness-of-fit plots generating using R (Version 2.13) based on the conditional weighted residuals distribution and the predicted versus observed tumor size measurements at both the population and individual levels. The tumor size model was developed to evaluate data from both treatment arms individually as well as simultaneously on the combined dataset.

2.5 Modeling Matched Cases from ECOG 4599 and CALGB 30203

The CALGB trial based parameter estimates for the linear growth rate and the treatment-related shrinkage rate differed from those originally published from the large FDA sample. To determine whether the deviation of the parameter estimates were specific to the CALGB data collection we extracted longitudinal tumor measurement data from a clinical trial conducted by the Eastern Cooperative Oncology Group (ECOG), trial 4599. That study compared front-line therapy with carboplatin and paclitaxel with or without the addition of bevacizumab (Sandler et al., 2006). The ECOG4599 data constituted more than one fourth the total sample used to generate the FDA model. One hundred three individual cases were selected from the carboplatin, paclitaxel, and placebo
arm of ECOG 4599 by matching to the CALGB 30203 sample on patient visit time, number of lesions, and sum of the longest diameter of the target lesions at baseline.

2.6 Calculation of time to tumor growth

In addition to change in tumor size at 8 weeks an alternative metric of treatment effects on serial tumor measurements, time to tumor growth (TTG) was evaluated (Claret et al., 2013) (Figure 4.3). TTG is expressed in time units (week) and depends on all the parameters of the empirical tumor model including baseline (Base, cm), tumor shrinkage rate constant (SR, 1/week) and tumor progression rate constant (PR, cm/week) (Eq 2).

\[
\frac{dT(TTG)}{dt} = Base \cdot e^{-SR(TTG)} \cdot (-SR) + PR = 0
\]
Figure 4.3. A **graphical representation of tumor matrices**.
TTG: time-to-tumor growth estimated using the FDA model; PTR: percentage tumor reduction from baseline.
2.7 Survival Analysis

In order to determine the relationship between tumor size change and overall survival, the tumor size model was used to predict tumor size change at early time points (4, 6 and 8 weeks). In addition to the predicted tumor size changes at 4, 6 and 8 weeks, time to tumor growth and other clinical risk factors (Table 4.2) were tested in Cox model for overall survival using pre-selection with a stepwise significance at 0.05. For the purpose of evaluating the prediction of time to death using measures of tumor dynamics, a parametric survival was adapted to estimate this relationship. Parametric survival functions including: exponential; Weibull; and log normal were tested, and the final survival distribution selected based on likelihood ratio test and Akaike information criterion.
### Table 4.2. Demographic and clinical characteristics of patients in CALGB 30203 and 30303 trial.

<table>
<thead>
<tr>
<th></th>
<th>CALGB30203 (n=103)</th>
<th>CALGB30303 (n=124)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>ECOG PS</strong></td>
<td>0 (n=40)</td>
<td>0 (n=61)</td>
</tr>
<tr>
<td></td>
<td>1 (n=58)</td>
<td>1 (n=63)</td>
</tr>
<tr>
<td></td>
<td>2 (n=5)</td>
<td></td>
</tr>
<tr>
<td><strong>SEX</strong></td>
<td>Male (n=63)</td>
<td>Male (n=82)</td>
</tr>
<tr>
<td></td>
<td>Female (n=40)</td>
<td>Female (n=42)</td>
</tr>
<tr>
<td><strong>AGE</strong></td>
<td>Median=61 (Range:25-83)</td>
<td>Median=61 (Range:29-78)</td>
</tr>
<tr>
<td><strong>Height</strong></td>
<td>Median=171 (Range:148-200)</td>
<td>Median=171 (Range:150-198)</td>
</tr>
<tr>
<td><strong>Weight</strong></td>
<td>Median=75 (Range:45-138)</td>
<td>Median=75 (Range:43-128)</td>
</tr>
<tr>
<td><strong>Histology</strong></td>
<td>Non-small cell (n=3)</td>
<td>Non-small cell (n=7)</td>
</tr>
<tr>
<td></td>
<td>Adenocarcinomas (n=50)</td>
<td>Adenocarcinomas (n=57)</td>
</tr>
<tr>
<td></td>
<td>Squamous (n=24)</td>
<td>Squamous (n=29)</td>
</tr>
<tr>
<td></td>
<td>Undifferentiated Large Cell (n=5)</td>
<td>Undifferentiated Large Cell (n=2)</td>
</tr>
<tr>
<td></td>
<td>Bronchoalveolar (n=4)</td>
<td>Bronchoalveolar (n=2)</td>
</tr>
<tr>
<td></td>
<td>Undifferentiated non-small cell (n=17)</td>
<td>Undifferentiated non-small cell (n=27)</td>
</tr>
<tr>
<td><strong>Ethnicity</strong></td>
<td>Hispanic (n=1)</td>
<td>Hispanic (n=0)</td>
</tr>
<tr>
<td></td>
<td>Non-Hispanic (n=89)</td>
<td>Non-Hispanic (n=115)</td>
</tr>
<tr>
<td></td>
<td>Unknown (n=13)</td>
<td>Unknown (n=9)</td>
</tr>
<tr>
<td><strong>Race</strong></td>
<td>White (n=85)</td>
<td>White (n=105)</td>
</tr>
<tr>
<td></td>
<td>Black (n=13)</td>
<td>Black (n=18)</td>
</tr>
<tr>
<td></td>
<td>Asian (n=2)</td>
<td>Asian (n=0)</td>
</tr>
<tr>
<td></td>
<td>Native Hawaiian or Pacific Islander (n=1)</td>
<td>Native Hawaiian or Pacific Islander (n=1)</td>
</tr>
<tr>
<td></td>
<td>American Indian or Alaska Native (n=1)</td>
<td>American Indian or Alaska Native (n=0)</td>
</tr>
<tr>
<td></td>
<td>Unknown (n=1)</td>
<td>Unknown (n=0)</td>
</tr>
<tr>
<td><strong>Prior_Chemo</strong></td>
<td>No (n=84)</td>
<td>No (n=124)</td>
</tr>
<tr>
<td></td>
<td>Yes (n=17)</td>
<td>Yes (n=0)</td>
</tr>
<tr>
<td></td>
<td>Missing (n=2)</td>
<td>Missing (n=0)</td>
</tr>
<tr>
<td><strong>Prior_RT</strong></td>
<td>No (n=102)</td>
<td>No (n=109)</td>
</tr>
<tr>
<td></td>
<td>Yes (n=0)</td>
<td>Yes (n=15)</td>
</tr>
<tr>
<td></td>
<td>Missing (n=1)</td>
<td>Missing (n=0)</td>
</tr>
<tr>
<td><strong>Disease stage</strong></td>
<td>IIIB (n=9)</td>
<td>IIIB (n=12)</td>
</tr>
<tr>
<td></td>
<td>IV (n=88)</td>
<td>IV (n=112)</td>
</tr>
<tr>
<td></td>
<td>Recurrent (n=6)</td>
<td>Recurrent (n=0)</td>
</tr>
</tbody>
</table>

Chemo: Chemotherapy; RT: Radiation.
2.8 Script to “Clean CRF Data” to Match RECIST 1.1 Consistently

When considered a conservative evaluation of longest diameter of tumor size, RECIST criteria 1.1 has been proposed to apply for trajectory selection. The tumor dynamic data was further manipulated by a script based program implemented using R (version 2.13) to extract and calculate the sum of longest diameter of the tumor size at each measurement using RECIST 1.1 criteria. The majority of tumors were measured by computed tomography scan (93.3%) and the rest of 6.62% were measured in different methods. Besides removing lesions with missing information and deleting individuals with baseline measurements only, the script based extraction was further applied to comprise the following steps: 1) exclude the lesions with inconsistent follow-up, 2) select up to a total of five lesions with a maximum of two lesions per organ and exclude lymph nodes if the maximum number of target lesions are exceeded.
R script code

#### import data #######

library(MIfuns)
library(lattice)
library(reshape)

d <-
read.csv(file="C:/Claire/NSCLC1/NSCLC30303TEST2.csv",skip=0,header=TRUE,sep="","",as.is=TRUE)
names(d) <-
c("ID","OID","VISIT","TREATMENT","LESION","LOCATION","SIZE","OLOCATION")

d$SIZE <- as.numeric(as.character(d$SIZE))

origi.dat <- d
write.table(d,file="C:/Claire/NSCLC1/30303TEST1.csv",sep="","",append=F,quote=F,col.names=T,row.names=F,na=".")

######## calculate sum of baseline tumor sizes #######

d1 <-  d[d$VISIT==0,]
reapply <- function(x,INDEX,FUN,...){
    y <- tapply(x,INDEX)
    z <- tapply(x,INDEX,FUN,...)
    z[y]
}

d1 <- transform(d1, basesum = reapply(SIZE, INDEX = ID, FUN = sum))
d1 <- d1[, c("ID", "basesum")]
d2 <- d1[!duplicated(d1$ID),]
d3 <- merge(d, d2, by.x = "ID", by.y = "ID", all.x = T)
write.table(d, file = "C:/Claire/NSCLC1/30303TEST2.csv", sep = ",", append = F, quote = F, col.names = T, row.names = F, na = ".")

######## remove lesion and size which are missing ########
d <- d3
d <- d[!is.na(d$LESION) & !is.na(d$SIZE),]
head(d)
write.table(d, file = "C:/Claire/NSCLC1/30303TEST3.csv", sep = ",", append = F, quote = F, col.names = T, row.names = F, na = ".")

###### determine which lesions are present for each individual at every visit ######

#### how many follow up+baseline in each patient
ni <- length(id <- unique(d$ID))
keep.lesion <- NULL
temp <- NULL
for (i in 1:ni){
    visit <- unique(d$VISIT[d$ID == id[i]])
nvisit <- length(visit)
s.lesion <- d$LESION[d$ID == id[i]]
u.lesion <- unique(s.lesion)
n.sl <- length(s.lesion)
n.ul <- length(u.lesion)

l <- data.frame(u.lesion,rep(0,n.ul))

names(l) <- c("lesion","times")

for (j in 1:n.ul) {
    for (k in 1:n.sl) {
        if (u.lesion[j]==s.lesion[k])
            l$times[j] <- l$times[j]+1
    }
}

l <- l[l$times==nvisit,]

temp <- data.frame(rep(id[i],length(l$lesion)),l$lesion,rep(nvisit,length(l$lesion))

keep.lesion <- rbind.data.frame(keep.lesion,temp)
}

names(keep.lesion) <- c("ID","LESION","NVISIT")

write.table(keep.lesion,file="C:/Claire/NSCLC1/30303TEST4.csv",sep="",append=F,quote=F,col.names=T,row.names=F,na=".")

######## delete IDs with baseline visit only ########

dd <- keep.lesion

dd <- dd[dd$NVISIT >= 2,]

######## calculate number of lesions that appear at all of the visits ########

ni <- length(dd$ID)

dd$flag <- rep(0,ni)
n <- ni-1

for (i in 1:n) {
    if (dd$ID[i]==dd$ID[i+1])
        dd$flag[i] <- 1
    else
        dd$flag[i] <- 0
}

dd <- transform( dd, NLESION=reapply(flag,INDEX=ID,FUN=sum)+1)

dd$flag <- NULL

keep.lesion <- dd

################ extract data with only the lesions that appear at all of the visits

a <- merge(d,keep.lesion,by=c("ID","LESION"),all.y=T)

a <- a[order(a$ID,a$VISIT,a$LESION),]

a <-

a[,c("ID","VISIT","LESION","SIZE","basesum","NLESION","TREATMENT","LOCATION","NVISIT")]

all.lesions <- a

write.table(all.lesions,file="C:/Claire/NSCLC1/30303TEST5.csv",
sep="",append=F,quote=F,col.names=T,row.names=F,na=".")

### add flag for lesions which are labeled as "LYMPH NODE" #######

flag <- rep(0,length(a$ID))

for (i in grep("LYMPH NODE",a$LOCATION)){

flag[i] <- 1
}
a$LYM <- flag
lym <- a[a$LYM==1,c("ID","LESION","LYM")]
lym <- unique(lym)
write.table(lym,file="C:/Claire/NSCLC1/30303TEST6.csv", sep="","append=F,quote=F,col.names=T,row.names=F,na=".")
a$LYM <- NULL
a <- merge(a,lym,by=c("ID","LESION"),all.x=T)
a <- a[order(a$ID,a$VISIT,a$LESION),]
a$LYM[is.na(a$LYM)] <- 0

write.table(a,file="C:/Claire/NSCLC1/30303TEST7.csv", sep="","append=F,quote=F,col.names=T,row.names=F,na=".")

######### calculate sum of lesions size at each visit for IDs with <= 3 lesions present at each visit #########
a1 <- a[a$NLESION <=3,]
aa1 <- aggregate(x=list(visitsum=a1$SIZE),by=list(ID=a1$ID,VISIT=a1$VISIT),FUN=sum)
aa1 <- aa1[order(aa1$ID,aa1$VISIT),]
a1 <- merge(a1,aa1,by=c("ID","VISIT"),all.x=T)
a1 <- a1[order(a1$ID,a1$VISIT,a1$LYM,-a1$SIZE),]
write.table(a1, file="C:/Claire/NSCLC1/30303TEST8.csv", sep="","append=F,quote=F,col.names=T,row.names=F,na=".")

######## IDs lesion # >3 ########

a2 <- a[a$NLESION >3,]

a2 <- a2[order(a2$ID,a2$VISIT,a2$LYM,-a2$SIZE),]

write.table(a2, file="C:/Claire/NSCLC1/30303TEST9.csv", sep="","append=F,quote=F,col.names=T,row.names=F,na=".")

######## identify three lesion numbers with biggest size at baseline and exclude "lymph node" for #lesion >=4 ########

d <- a2[a2$VISIT==0,c("ID","LESION")]

d1 <- NULL
temp <- NULL

ni <- length(id<-unique(d$ID))

for (i in 1:ni){
  lesion <- unique(d$LESION[d$ID==id[i]])
  nlesion <- length(lesion)
  temp <- data.frame(rep(id[i],3),lesion[1:3])
  d1 <- rbind.data.frame(d1,temp)
}

names(d1) <- c("ID","LESION")

d1 <- d1[order(d1$ID,d1$LESION),]

v <- d1

############## calculated sum of three largest lesion sizes at each visit ##############
v1 <- merge(v,a2,by=c("ID","LESION"),all.x=T)

vv1 <-
aggregate(x=list(visitsum=v1$SIZE),by=list(ID=v1$ID,VISIT=v1$VISIT),FUN=sum)
v1 <- merge(v1,vv1,by=c("ID","VISIT"),all.x=T)
a2 <- v1

a2 <- a2[order(a2$ID,a2$VISIT,a2$LYM,-a2$SIZE),]
a2$NLESION <- 3

########## combine two subset together #######
a <- rbind.data.frame(a1,a2)

###a$TREAT <- ifelse(a$TREATMENT=='PLACEBO',0,1)###
a <- a[,c("ID","VISIT","LESION","SIZE","visitsum","basesum","NLESION"
,"LYM","TREATMENT","NVISIT")]
names(a) <- c("ID","VISIT","LESION","SIZE","VSUM","BSUM","NLESION"
,"LYM","TREATMENT","NVISIT")
a <- a[order(a$ID,a$VISIT,a$LYM,-a$SIZE),]

######## export dataset with all ids ###########
write.table(a,file="C:/Claire/NSCLC1/NSCLC30303FINAL.csv",
sep="",append=F,quote=F,col.names=T,row.names=F,na=".")
2.9 Blinded Re-Evaluation of Imaging Data

To identify sources of variance between patient outcomes and the modeled change in tumor size over time, we obtained the original sets of images from patients enrolled at the University of Chicago in studies 30203 and 30303. One radiologist, blinded to the original case report forms and radiology reports (co-author AO) reviewed all of the baseline images and identified and measured all target lesions. Unblinded to the lesions, but still blinded to the original measurements, the radiologist performed serial measurements on the identified target lesions on the University of Chicago patients.

3. Results

3.1 Data Quality Control

We discovered that 9730 study case report forms (CRFs) frequently included no change (nc) or not available (na) rather than tumor size measurements on subsequent CT scans as in Figure 4.2. Consequently the entire trial data set had too much missing data to be useful for validating the longitudinal tumor growth model and data from all 561 subjects was excluded.

For the CALGB trials, we subjected the patients’ data to the same standard for inclusion as in the FDA model (at least a baseline measurement and measurements recorded at some subsequent time-point). Figure 4.1 describes the attrition for 140 original cases in the CALGB 30203 trial and 160 in CALGB 30303. This resulted in a total of 227 patients available for the analyses.

3.2 Longitudinal Modeling of Tumor Size

Parameter estimates for sum of longest tumor dimensions at baseline (M_BASE),
the treatment-effect/shrinkage rate (M_SR), and the linear tumor growth rate (M_PR) were determined and compared to the results of similar study arms from the original study as shown in Table 4.3. The individual predictions vs. observations of sum of the longest diameter were compared in each individual between FOCE and SAEM methods. The final parameters were estimated using SAEM based on individual fitting.
<table>
<thead>
<tr>
<th>Study</th>
<th>Treatment</th>
<th>N</th>
<th>M_BASE (cm)</th>
<th>M_SR (1/week)</th>
<th>M_PR (cm/week)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDA trial treatment arms (E4599 &amp; TAX 326)</td>
<td>PCB</td>
<td>434</td>
<td>9.1(0.33)</td>
<td>0.06(0.004)</td>
<td>0.13(0.02)</td>
</tr>
<tr>
<td></td>
<td>PC</td>
<td>444</td>
<td>8.0(0.30)</td>
<td>0.038(0.01)</td>
<td>0.14(0.04)</td>
</tr>
<tr>
<td></td>
<td>DC</td>
<td>408</td>
<td>8.7(0.31)</td>
<td>0.052(0.01)</td>
<td>0.16(0.02)</td>
</tr>
<tr>
<td></td>
<td>DCb</td>
<td>406</td>
<td>9.2(0.38)</td>
<td>0.047(0.005)</td>
<td>0.16(0.02)</td>
</tr>
<tr>
<td></td>
<td>VC</td>
<td>404</td>
<td>8.5(0.28)</td>
<td>0.063(0.01)</td>
<td>0.17(0.02)</td>
</tr>
<tr>
<td>NCI trial treatment arms (CALGB30203)</td>
<td>GCb+/- Zi or Ce</td>
<td>103</td>
<td>8.14(0.53)</td>
<td>0.014(0.005)</td>
<td>0.041(0.016)</td>
</tr>
<tr>
<td>(CALGB30303)</td>
<td>DC+/- BNP</td>
<td>124</td>
<td>8.66(0.48)</td>
<td>0.037(0.005)</td>
<td>0.079(0.011)</td>
</tr>
<tr>
<td>(CALGB30203+CALGB30303)</td>
<td>GCb+/- Zi or Ce and DC+/- BNP</td>
<td>227</td>
<td>8.44(0.35)</td>
<td>0.027(0.003)</td>
<td>0.066(0.008)</td>
</tr>
</tbody>
</table>

PCB, Paclitaxel, Carboplatin and Bevacizumab; PC, Paclitaxel and Carboplatin; DC, Docetaxel and Cisplatin; DCb, Docetaxel and Carboplatin; VC, Vinorelbine and Cisplatin; GCb, Carboplatin and Gemcitabine; Zi, Zileuton; Ce, Celecoxib; BNP, BNP7787

Table 4.3. **Tumor model parameter estimates and precision standard error (SE) of baseline (M_BASE), shrinkage rate (M_SR) and progression rate (M_PR) for FDA registration trials and CALGB 30203 and 30303 trial.**
Variance in parameter estimates increased as sample size was reduced from typical phase III to typical phase II size study arms. With a combination of both 30203 and 30303 trials, the model estimates of baseline tumor size, shrinkage rate and progression rate were 8.44 cm, 0.027 1/week and 0.066 cm/week, respectively. For example a patient with an average baseline tumor size at 8.1 cm will after 1 month, have the typical tumor burden decrease to $8.44\times e^{(-0.027 \times 4)}+ 0.066 \times 4=7.84$ cm. This 7.1% decrease reflects the drug effect on tumor size. Compared to the average parameter estimates from five treatment arms of FDA data, the CALGB results underestimated $M_{\text{BASE}}$, $M_{\text{SR}}$, $M_{\text{PR}}$ in 3%, 48% and 57%, respectively.

These differences in tumor size and growth metrics between the smaller CALGB dataset and the FDA dataset were associated with tumor size at 8 weeks no longer being a statistically significant predictor of survival in NSCLC (Table 4.4). This inconsistency in the prediction of survival by tumor size with data collected by an experienced cooperative clinical trials group undermined plans to replace RECIST with continuous measures of tumor burden in smaller, prospective, randomized phase II trials. We conducted additional studies to identify factors in acquisition of these data that might have led to this unexpected finding.

Furthermore, no significant changes in the tumor parameter estimates were observed when applying additional script-based filtration by adding the restriction of number of the lesions, consistence of follow-up and number of lymph nodes.
Table 4.4. **Comparison of tumor parameter estimates between CALGB30203 trial and a subset of E4599 trial which matched to CALGB30203 tumor information.**

<table>
<thead>
<tr>
<th>Study</th>
<th>Treatment</th>
<th>N</th>
<th>M_BASE (cm)</th>
<th>M_SR (1/week)</th>
<th>M_PR (cm/week)</th>
</tr>
</thead>
<tbody>
<tr>
<td>FDA trial treatment arms</td>
<td>PC</td>
<td>444</td>
<td>8.0</td>
<td>0.038</td>
<td>0.14</td>
</tr>
<tr>
<td>(E4599&amp; TAX 326)</td>
<td>PC subgroup</td>
<td>103</td>
<td>9.26</td>
<td>0.0138</td>
<td>0.0346</td>
</tr>
<tr>
<td>NCI trial treatment arm (CALGB30203)</td>
<td>GCb +/- ZI or Ce</td>
<td>103</td>
<td>8.1</td>
<td>0.0139</td>
<td>0.0407</td>
</tr>
</tbody>
</table>

PC, Paclitaxel and Carboplatin; GCb, Carboplatin and Gemcitabine; ZI, Zileuton; Ce, Celecoxib
3.3 Evaluation of deviations in parameter estimates and survival prediction

First we hypothesized that these lower estimates might reflect lower quality of data from small cooperative group trials compared to perhaps more meticulously accurate data submitted to the FDA for review. We therefore matched 103 patients from the ECOG 4599 study carboplatin/paclitaxel arm, part of the dataset used to generate the FDA model, with the 103 CALGB 30203 cases (who received carboplatin/gemcitabine). For this subset of matched cases in ECOG 4599 the mean baseline tumor size was larger, but the parameter estimates for M_SR and M_PR were now more similar to CALGB 30203 than to the results for the entire 444 subject carboplatin/paclitaxel arm of ECOG 4599 (Table 4.4). This implied that the deviation of parameter estimates between the CALGB data and the similar, but larger, treatment arms in the FDA dataset were unlikely to be due to significant differences in data quality between trials conducted by different groups, and instead reflected effects of decreasing the size of the analyzed subject pool.

Some recent studies suggested that TTG might be more robust than tumor size at early fixed time-points for colorectal cancer (Claret et al., 2013) and NSCLC (Claret et al., 2014). We therefore tested whether TTG might be a significant predictor of survival in the CALGB 30203 and the matched E4599 smaller dataset. In the Cox proportional hazard model for CALGB 30203 patients, the baseline tumor size (centered at 8.5cm) (P<0.001) and TTG (P<0.001) were significant predictors of time to death when incorporated as independent factors. The p-value of each coefficient was no longer significant when both factors were tested in the survival model simultaneously. The TTG and baseline tumor size were correlated (r = 0.75) (P<0.001). Therefore, the final survival model included the baseline tumor size and ECOG performance status (0/1/2) (Eq 3).
log(T)=\beta_0 + \beta_1 \times (\text{baseline-8.5}) + \beta_2 \times \text{ECOG} + \varepsilon

where T is time to tumor death (day) in a log normal distribution, \( \beta_0 \) is the intercept when no covariate was added, \( \beta_1 \) and \( \beta_2 \) are the coefficients for baseline-8.5 and ECOG, respectively, and \( \varepsilon \) represents the residual error. In the matched subset of E4599 patients, TTG remained a statistically significant predictor in this model. However, when tested in the remaining 232 subjects of the carboplatin/paclitaxel arm, TTG was no longer a significant predictor of survival. We concluded that this loss of tumor size as a significant predictor of survival in smaller datasets was not an effect of the specific parameter of continuous measurement of tumor burden.

A difficult-to-test hypothesis is that the model accurately reflects that the CALGB 30203 and E4599 subsets of patients are genuinely different from the larger population of patients on which the FDA model was based. Our experience with the multi-step process of CT-imaging measurement and transmission of measurements into clinical trial databases offers an alternative hypothesis: the routine variance in tumor burden measurements introduced by our current RECIST-oriented clinical trial methods contributes significantly to this loss of power because of the inherent variance introduced with this approach. This variance further confounds computational models of continuous tumor growth reducing their ability to accurately predict survival in patient populations typical of phase 2 trials.

We therefore explored specific modifiable factors in the collection and reporting of tumor measurements might contribute to the altered shrinkage rate and growth rate parameter estimates in the longitudinal growth model when the size of the population was decreased. To evaluate the reproducibility of the tumor measurements on the case report
forms, an independent radiologist in blinded fashion re-determined and measured the baseline target lesions from the original CT images collected from 15 patients enrolled in CALGB 30203 and 30303 at one institution (Figure 4.4A). For 4 of the 15 patients, at least one additional target lesion was identified. Once unblinded to the determination of target lesions, the radiologist performed blinded serial measurements of the designated lesions (Figure 4.4B). Although displayed for consistency, 3 of the 15 subjects did not have an on-treatment assessment and therefore were not included in subsequent modeling analyses. For the 12 cases with serial measurements, 4 (subjects 7, 8, 9, 12) had trajectories of the measured sums of longest dimensions that were nearly superimposable between the case report forms (CRF) and the blinded evaluator re-assessment. Four (subjects 1, 3, 4, and 5) had obvious divergence between the CRF and blinded evaluations in terms of the magnitude of change in tumor burden and time-points at which these changes are registered. The remaining 4 had differences of unclear significance (2, 6, 10, 11).

3.5 Estimated impact of continuous measurement variance on modeled endpoints

RECIST was developed to be robust to inter-rater variance in measurements by setting categories for tumor size changes (Progressive Disease, Partial Response, and Complete Response) based on thresholds for magnitudes of change that would be unlikely to be due to the greatest degree of inter-rater variance (Therasse et al., 2000). A patient’s category of response would then likely only be due to a significant effect of treatment (Moertel et al., 1976; Miller et al., 1981). It is therefore not surprising that in settings where inter-rater variance is not actively controlled, assessments of continuous
measurements of tumor growth will not improve upon our current RECIST-based categorical and time-to-event strategies.

We hypothesized that this inter-rater variance in tumor burden assessments would have a significant effect on a continuous-measurement-based strategy such as TTG with less effect on a RECIST-based time to event endpoint such as progression-free survival (PFS). For the 12 subjects with serial CRF and blinded radiologist measurements (Table 4.5), we identified an average 25% delay in TTG calculated from the re-evaluated scans compared with the CRF data, but no differences in PFS assessments. Despite differences in target lesion assessment and measurement, subjects met criteria for progressive disease at the same imaging session in both datasets.
Figure 4.4. **Comparisons of identified target lesion measurements from original case report forms and by blinded radiologist reviewer.**

A) Target lesion measurements from CALGB 30203 and 30303. X-axis; Yellow circles; CRFs (case report forms), Blue circles; BE (Blinded re-evaluation of imaging data by radiologist from the University of Chicago). Y-axis; Each circle depicts one lesion at baseline (BL) visit in centimeters (cm). Gray line between yellow and blue circle connects the same lesion.

B) Sum of the longest dimensions (cm) at each assessment time point for subjects in A over the course of the trial.
Table 4.5. **Comparisons of PFS and calculated TTG from the target lesion measurements by blinded radiologist evaluation (BE) and case report form (CRF) data**

<table>
<thead>
<tr>
<th>Patient ID</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PFS</strong> (week)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CRF</td>
<td>8</td>
<td>26</td>
<td>4</td>
<td>48</td>
<td>28</td>
<td>128</td>
<td>40</td>
<td>12</td>
<td>36</td>
<td>48</td>
<td>24</td>
<td>18</td>
</tr>
<tr>
<td>BE</td>
<td>8</td>
<td>26</td>
<td>4</td>
<td>48</td>
<td>28</td>
<td>128</td>
<td>40</td>
<td>12</td>
<td>36</td>
<td>48</td>
<td>24</td>
<td>18</td>
</tr>
<tr>
<td><strong>TTG</strong> (week)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CRF</td>
<td>1</td>
<td>20</td>
<td>74</td>
<td>27</td>
<td>43</td>
<td>53</td>
<td>21</td>
<td>43</td>
<td>23</td>
<td>20</td>
<td>13</td>
<td>46</td>
</tr>
<tr>
<td>BE</td>
<td>51</td>
<td>31</td>
<td>94</td>
<td>36</td>
<td>52</td>
<td>73</td>
<td>22</td>
<td>52</td>
<td>24</td>
<td>13</td>
<td>26</td>
<td>47</td>
</tr>
</tbody>
</table>
4. Discussion

This examination of a NSCLC longitudinal tumor growth model and endpoints in published CALGB studies revealed current limitations to employing continuous measurements of tumor burden in phase II clinical trials. Modeling of typical phase III clinical trial-size samples has reproducibly demonstrated tumor burden metrics as predictors of survival (Claret et al., 2012; Stein et al., 2011; Claret et al., 2013; Stein et al., 2012). Contrary to our expectations, we showed that decreasing the sample size to that of a typical phase II trial results in complete loss of the predictive power of these tumor metrics. Even though TS at 8 weeks is no longer a statistically significant predictor of survival in NSCLC with a typical phase II sample size, the direction of this predictor to survival remains the same as observed in the large phase III trials. As Bonate illustrated in a simulation study based on the FDA model structure, the parameter bias did not translate to prediction bias in tumor size as the measurement error with the expected bounds for tumor size assessment (Bonate et al., 2013). This suggests that we cannot rule out the clinical utility of TS at early time points in phase II survival prediction because of the low predictive power in a small size trial. In fact, other sources of variance such as phase II survival data in contributing to the model predictability may also need to be evaluated. Closer scrutiny of the original images and the recorded data revealed variance in the process by which tumor burden is assessed and recorded to meet RECIST standards. This variance has no apparent effect on RECIST categories or time-to-event endpoints, but does affect tumor burden metrics.

There is no superior alternative approach to RECIST for the standardized assessment of anatomical tumor burden and its change over time (Eisenhauer et al., 2009;
This categorical system applied to quantitative assessments of tumor burden provides low inter-rater variance (progressive disease will be determined with high uniformity across sites in a multi-center trial and among trials) at the expense of efficiency (requires more patients to be observed over long periods of time). Our findings are consistent with investigators collecting and curating the quantitative tumor burden data with sufficient precision to support use of RECIST but not to support more computationally intensive methods of evaluating effects of treatments in small clinical trials. It is no surprise that many investigators have found no significant advantages to use of quantitative methods (such as tumor size ratio) over more qualitative time-to-event strategies (such as progression-free-survival) for predicting impact on overall survival (Fridlyand et al., 2011; An et al., 2011; Kaiser et al., 2013).

This study had a limited data sample for analysis, but it required significant effort to obtain these data. The qualitative documentation of tumor response data in earlier trials was not suitable for further quantitative analysis. For the more recent trials, the primary databases maintained the RECIST-based categories in data fields, but obtaining the quantitative tumor measurements required manual retrieval and processing of archived paper forms. The small cohort of patients for whom images were available and reviewed might have been a biased sample, but this site had been a major contributor to enrollment across thoracic oncology trials in CALGB with the stringent audit and quality control processes applied for member sites. The data are therefore likely representative of the overall quality of data in the larger clinical trials. Furthermore, an independent trial, conducted by a completely different set of institutions (ECOG) yielded similar results. We cannot exclude the possibility that this particular subset of patients matched between
the CALGB and ECOG datasets represents a unique group of NSCLC patients whose
tumor growth patterns are distinct from the typical patient population. Therefore our
findings will require confirmation in other datasets.

The study demonstrates a potential major flaw in assumptions on the use of
quantitative measures of tumor burden and computational analysis. The process of
measuring, transmitting, analyzing, and interpreting CT imaging-based measures of
tumor contributes significant, but potentially modifiable variance. Centralized collection
and measurement of CT images with semi-automated and digitally enhanced procedures
should significantly reduce this variance. Advances in computing and digital data
management in the past several years have reduced costs and make paperless systems
with fewer opportunities for manual error possible. Our findings suggest that
establishing methods with less inter-rater variance should be a worthwhile investment in
the future of cancer therapeutics assessment.
CHAPTER V: Clinical Trial Simulation to Evaluate Study Design: Capturing Abiraterone and Nilotinib Exposures

1. Introduction

In the past twenty years, oral anticancer drug therapy has become more prevalent (Singh et al., 2004). Unlike intravenous drug administration, orally administered agents undergo absorption from the gastrointestinal lumen through the intestinal epithelium into the portal vein and then into the systemic circulation. The fraction of an orally administered drug dose that reaches the systemic circulation is the drug’s oral bioavailability and significant differences or changes in bioavailability will lead to a significant variation in drug exposure (Martinez et al., 2002) and thus modified therapeutic and or toxic effects.

One of most well-known factors contributing to variability in oral bioavailability is food intake (Singh et al., 2004; Winstanley et al., 1989; Gu et al., 2007). The effect of food on oral bioavailability has the potential to cause a clinically significant impact by causing variable systemic drug exposures that can lead to drug toxicity or therapeutic failure (Koch et al., 2009; Kang et al., 2010). Furthermore, the alternation of pharmacokinetic profile of several standard and investigational anticancer drugs such as busulfan, topotecan and fluorouracil caused by food-drug interactions have been previously reported (Singh et al., 2004). Quantifying the effect of food on drug exposure is important when designing a clinical trial for oral anti-cancer drugs. In addition to food related variability, variability in drug pharmacokinetic parameters between individuals as well as between occasions within an individual also contributes to variability in drug exposure. In order to assess the contribution of different magnitudes of between-
individual and between-occasion variability on drug exposure, this study used a population PK modeling approach and Monte Carlo methods to simulate a virtual clinical trial of patients who took drug in both the fasted and fed states.

Such a clinical trial simulation framework provides insight as to whether a particular study design, with a random sampling schema to reflect a typical clinical practice setting, permits the retrieval of pharmacokinetic (PK) parameters and their variability under different prandial conditions. In addition, simulation of a clinical trial can assess how patient specific covariates, between-individual, and between-occasion variability affect the ability to accurately capture individual level exposures. The aim of this study was to evaluate whether a pre-defined oncology trial with a sparse drug concentration sampling schedule can adequately capture individual level drug exposures, random variability and the food effects of (1) abiraterone (92% decrease in oral clearance), and (2) nilotinib, (18% decrease in oral clearance). To achieve these objectives, a virtual cancer study population was simulated with assigned pharmacokinetic characteristics, food intake and between-individual as well as between-occasion variability. A population pharmacokinetic (Pop PK) approach was utilized to retrieve the Pop PK parameters under these conditions and examine whether or not these parameters could be adequately retrieved on estimation at both the individual and population level.

2. Materials and Methods

2.1 Simulation of Patient Pharmacokinetic Characteristics

Hypothetical patients were created with fasting state Pop PK characteristics mimicking those found in oncology patients. (See Table 5.1 for abiraterone (Ryan et al.,
2010) and Table 5.2 for nilotinib (ZYTIGA, Tanaka et al., 2010) patient characteristics. The Pop PK parameters were assumed to be log-normally distributed and the parameter values for simulation were obtained or calculated based on literature values from a non-compartmental analyses (Ryan et al., 2010; ZYTIGA, Tanaka et al., 2010). A one compartment model with first order absorption and elimination was assumed as the model structure. Between-individual variability of oral clearance (CL/F) and volume of distribution (Vd/F) were based on the mean and standard deviation of pharmacokinetic parameters from published studies. Abiraterone between-individual variability values for oral clearance and volume of distribution were 174% and 73%, respectively, in the fasted state and 42.2% and 36%, respectively, in the fed state [7]. The between-individual variability of the absorption rate constant was assumed to be 20% and a proportional residual error with 30% CV was assumed based on literature values for abiraterone (Ryan et al., 2010). For nilotinib, the between-individual variability of oral clearance and volume of distribution values were set to 55% and 37% CV, respectively, for both fed and fasting states (Tanaka et al., 2010). A between-individual variability for the nilotinib absorption rate constant of 20% and a proportional residual error with 10% CV based on the lower limit of quantification of the nilotinib concentration assay were assumed (Tanaka et al., 2010).

Values of the Pop PK parameters for each hypothetical patient were randomly chosen from their respective distributions with a correlation of 0.6 applied between clearance and volume.
Table 5.1. **Population pharmacokinetic parameters for abiraterone patients**

<table>
<thead>
<tr>
<th>Abiraterone</th>
<th>Fasting status</th>
<th>Fed status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Parameter</td>
<td>Mean (SD)</td>
<td>Mean (SD)</td>
</tr>
<tr>
<td>Apparent Clearance / bioavailability [CL/F] (L/h)</td>
<td>2650 (4617)</td>
<td>231 (97.7)</td>
</tr>
<tr>
<td>Apparent Volume of distribution / bioavailability [Vd/F] (L)</td>
<td>25494 (18670)</td>
<td>4069 (1462)</td>
</tr>
<tr>
<td>Absorption rate (1/h)</td>
<td>1.65 (0.33)</td>
<td>1.65 (0.33)</td>
</tr>
</tbody>
</table>

F = oral bioavailability
Table 5.2. **Population pharmacokinetic parameters for nilotinib patients**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Fasting status</th>
<th>Fed status</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean (SD)</td>
<td>Mean (SD)</td>
</tr>
<tr>
<td>Apparent Clearance / bioavailability <a href="L/h">CL/F</a></td>
<td>33 (18)</td>
<td>27(15)</td>
</tr>
<tr>
<td>Apparent Volume of distribution / bioavailability [Vd/F] (L)</td>
<td>720 (267)</td>
<td>604(181)</td>
</tr>
<tr>
<td>Absorption rate (1/h)</td>
<td>0.74 (0.15)</td>
<td>0.74 (0.15)</td>
</tr>
</tbody>
</table>

F = oral bioavailability
2.2 Food Effects

For both abiraterone and nilotinib, it was assumed that 50% of doses were taken with food and that the values of the oral clearance and volume of distribution depended upon whether a particular dose was taken with food or without food.

To mimic the actual drug administration protocol for food intake, we considered a different food effect in the abiraterone versus nilotinib trials. This was done to test whether the model would be able to capture two extreme food effects, the between-individual and between-occasion variability, and the individual level exposure. For abiraterone, patients in the fed prandial state were assumed to have had a high fat meal given the availability of Pop PK parameter values in the reference for simulation from this state; therefore, the oral clearance and volume of distribution were reduced by 92% and 85%, respectively, if the dose was taken with food (Ryan et al., 2010). In contrast to the abiraterone simulations, patients who were simulated as receiving nilotinib were assumed to have a light fat meal and possibly just a glass of grape juice 2 hours before or 1 hour after nilotinib was taken. An 18% reduction in apparent clearance was introduced to reflect this food effect (Tanaka et al., 2010; Yin et al., 2010).

2.3 Study Design

Patients in the simulated abiraterone trials were randomly assigned to take 1,000 mg once daily and patients in the simulated nilotinib trials were assigned to take a 300 or 400 mg tablet twice daily in 1:1 fashion. Virtual pharmacokinetic samples for each patient were assumed to occur at week 1, week 4, and month 2 and month 3 based on a previously established trial sampling schedule (see Figure 5.1) after the start of the trial so each individual had a total of 4 measurements (one sample obtained on each occasion).
This sampling schedule was designed to fit in with the standard clinical follow up of the patient. The samples were assumed to occur at random times during clinic hours. The visit day for each pharmacokinetic sample of each hypothetical patient was randomly selected from a discrete distribution with 50% of virtual patients having a visit sample drawn on the recommended day, 20% ± 1 day, 20% at ± 2 days and 10% at ± 3 days from the scheduled visit date. Each clinic visit time was randomly selected from a uniform distribution assuming regular office hours from 7am to 6pm. To mimic food intake when patients take their medications, a random food covariate was generated for each hypothetical patient and sampled for the four different clinic visit occasions within each patient.

The hypothetical patients were assigned into trial sizes of 20, 50 or 70 individuals, and each trial was replicated 100 times. The same study design was tested using three different between-occasion variability levels: 10, 25, and 40% CV on drug clearance.
Figure 5.1. Study schema of abiraterone and nilotinib clinical trials.
2.4 Population Pharmacokinetic Model Estimation

The population pharmacokinetics of the hypothetical patients in each trial were analyzed using nonlinear mixed effect modeling software, NONMEM, Version VII (GloboMax_LLC, Ellicott City, MD, USA) using Wings for NONMEM, Version 7. One-compartment structural model with a proportional residual error model was tested using the first-order conditional estimation (FOCE) method. Between-individual variability was tested on oral clearance and volume of distribution, and between-occasion variability was tested on oral clearance only. The between-individual and between-occasion variability for Pop PK parameters were assumed to be log-normally distributed and evaluated using an exponential model $P_{i,k} = P_{TV} x e^{\eta_{p,i}} x e^{\eta_{p,k}}$ where $P_{i,k}$ is the parameter estimate for the $i^{th}$ individual after the $k^{th}$ dose administered and $P_{TV}$ is the typical value for the parameter at the population level. The variability between the $i^{th}$ individual and the population parameter value was described by $\eta_{p,i}$ which was identically distributed with a mean of 0 and a variance of $\omega_{\eta_{p,i}}^2$ (Feng et al., 2006). The variability between the $k^{th}$ occasion of dose administration and the population parameter value was described by $\eta_{p,k}$ which was identically distributed with a mean of 0 and a variance of $\omega_{\eta_{p,k}}^2$. In addition to the between-individual and between-occasion variability, residual variability was described by a proportional error model. That is, $y_{ij} = \hat{y}_{ij}(1 + \epsilon_{ij})$ where $y_{ij}$ and $\hat{y}_{ij}$ are the $j^{th}$ observed nilotinib or abiraterone concentration and its corresponding model predicted concentration, respectively, with the difference described by $\epsilon_{ij}$ which is assumed to be normally distributed with a mean of 0 and a variance of $\sigma^2$. In order to estimate between-occasion variability in the model, the residual error was fixed in the estimation step.
2.5 Evaluation of Clinical Trial Designs

The food effect in each trial was evaluated by comparing the fit of a base model with no food effects with a model that includes a food effect covariate on oral clearance. The model comparison was based on a likelihood ratio test using the objective function value (OFV) from NONMEM. The change in the OFV returned by NONMEM is approximately equal to $-2 \times \log \text{likelihood}$. The difference in $-2 \times \log \text{likelihood}$ between two models that are nested follows a $\chi^2$ distribution. The significance level for identifying the food effect corresponded to a decrease in the OFV of greater than 6.63 ($p\text{-value} \leq 0.01, \text{df}=1$). The power to detect a food effect is the percent of the trials where the population PK analysis demonstrated significant OFV change among the 100 trials in both abiraterone and nilotinib and retrieving a food effect value within 20% of the true food effect value.

In addition to determining whether or not a food effect was detected by the trial design, the accuracy and precision of the model for retrieving parameter values of clearance, between-individual and between-occasion variability was assessed using two statistical standard criteria: percent bias and percent precision (Huang et al., 2007).

At the population level, the bias of each parameter represents the difference between the estimated values from the simulated (true) value of the population, and the percent bias is the mean predicted error normalized by the simulated value taken from the literature. The bias is calculated using the equation shown below:

$$\text{Population \% bias} = \frac{1}{m} \sum_{j=1}^{m} \left( \frac{Y_{p,j} - Y_e}{Y_e} \right) x 100\%$$
where \( Y_{p,j} \) is the predicted value for the \( j^{th} \) trial in total \( m \) trials with the same given between-occasion variability level and the same sample size, and \( Y_e \) is the literature value we simulated from. The precision was estimated by calculating the root mean square error, reflecting the distribution of variance, and the percent precision is the root mean square error normalized by the simulated mean. This included the bias and precision of the estimation of the food effect.

\[
\text{Population \% precision} = \sqrt{\frac{\sum_{j=1}^{m} (\frac{Y_{p,j} - Y_e}{Y_e})^2}{m}} \times 100\%
\]

Parameter estimates at the individual level were evaluated using percent bias and percent precision.

\[
\text{Individual \% bias} = \frac{1}{n} \sum_{i=1}^{n} \left( \frac{Y_{p,i} - Y_{e,i}}{Y_{e,i}} \right) \times 100\%
\]

where \( Y_{p,i} \) and \( Y_{e,i} \) are the predicted and simulated values for the \( i^{th} \) patient, respectively. \( n \) represents the number of patients in the trial.

\[
\text{Individual \% precision} = \sqrt{\frac{\sum_{i=1}^{n} (\frac{Y_{p,i} - Y_{e,i}}{Y_{e,i}})^2}{n}} \times 100\%
\]

Then, an average of individual percent bias and precision of the \( m \) number of trials was calculated.

3. Results

The goodness-of-fit plots for the final model were shown in Figure 5.2 and 5.3 for abiraterone and nilotinib, respectively. The percent bias and precision for the population
oral clearance for abiraterone ranged from -37% to -31% and 36.2% to 38.1%, respectively (Table 5.3). The average percent bias and precision are noticeably smaller in magnitude for individual clearance estimates (range of 1.2% to 4.4% and 14.5% to 19.9%, respectively). This indicates that the model estimates for oral clearance are more accurate and less variable at the individual level compared to the population level. Across three between-occasion variability levels (10, 25, and 40%), 21%, 10% and 11% of the abiraterone trials, respectively, retrieved population clearance values within 20% of the true value. The ranges of percent bias and precision for between-individual variability were -45.1% to 42.1% and 42.8% to 45.9%, respectively, with minimal variations with number of patients in the trial or between-occasion variability. The ranges of percent bias and precision of between-occasion variability were -30% to -1.1% and 11.6% to 40.7%, respectively. There was a decrease in both the between-occasion variability bias and precision as the number of patients and between-occasion variability increased. The known food effect on oral clearance for abiraterone was identified in 100% of simulated trials with 20, 50 and 70 patients for the 10%, 25%, and 40% between-occasion variability levels. The ranges of percent bias and precision of food effect were 2.01% to 4.42% and 6.81% to 14.4%, respectively.
Figure 5.2. Goodness-of-fit plots for the final model of abiraterone.
Plot of population and individual predicted versus observed concentrations and plot of conditional weighted residuals versus population predicted concentrations and time (hours).
Figure 5.3. **Goodness-of-fit plots for the final model of nilotinib.**
Plot of population and individual predicted versus observed concentrations and plot of conditional weighted residuals versus population predicted concentrations and time (hours).
Table 5.3. Bias and precision of abiraterone oral clearance at the population level and individual level, between-individual variability, and between-occasion variability

<table>
<thead>
<tr>
<th>Between-occasion variability</th>
<th>Number of patients in trial</th>
<th>Oral Clearance – Population level</th>
<th>Oral Clearance – Individual level</th>
<th>Between-individual variability</th>
<th>Between-occasion variability</th>
<th>Food effect</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Bias</td>
<td>Precision</td>
<td>Bias</td>
<td>Precision</td>
<td>Bias</td>
</tr>
<tr>
<td>10%</td>
<td>20</td>
<td>-32.3%</td>
<td>37.1%</td>
<td>1.5%</td>
<td>14.5%</td>
<td>-44%</td>
</tr>
<tr>
<td></td>
<td>50</td>
<td>-36.6%</td>
<td>37.8%</td>
<td>1.3%</td>
<td>14.6%</td>
<td>-43.6%</td>
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<td></td>
<td>70</td>
<td>-36.8%</td>
<td>37.8%</td>
<td>1.4%</td>
<td>14.9%</td>
<td>-43.3%</td>
</tr>
<tr>
<td>25%</td>
<td>20</td>
<td>-31.7%</td>
<td>36.6%</td>
<td>3.3%</td>
<td>19.6%</td>
<td>-45.1%</td>
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<tr>
<td></td>
<td>50</td>
<td>-36.3%</td>
<td>37.6%</td>
<td>1.2%</td>
<td>18.9%</td>
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<td>36.2%</td>
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<td>70</td>
<td>-37%</td>
<td>38.1%</td>
<td>4.4%</td>
<td>19.6%</td>
<td>-42.6%</td>
</tr>
</tbody>
</table>
The percent bias and precision on the population oral clearance for nilotinib ranged from -13.3% to -11.8% and 14.2% to 17.0%, respectively (Table 5.4) and were consistent across between-occasion variability levels (10%, 25% and 40%). In contrast to abiraterone, the average individual nilotinib oral clearance estimates were significantly more accurate and precise than population estimates with the percent bias and precision ranging from -1.9% to -0.5% and 4.2% to 8.6%, respectively. Across three between-occasion variability levels (10, 25 and 40%), 86, 84 and 83% of the nilotinib trials, respectively, retrieved the population oral clearance within 20% of the true value. The ranges of percent bias and precision of between-individual variability were -9.9% to 7.9% and 11.3% to 19.4%, respectively and ranged from -3.9% to -0.4% and 4.9% to 11.0% for percent bias and precision on between-occasion variability. Retrieval of the known food effect in this system was observed in 100% of the simulated nilotinib trials with 10% between-occasion variability with trial sizes of 25, 50, and 70 patients. For nilotinib trials simulated with 25% between-occasion variability, significant food effects on oral clearance were retrieved in 80% of 20 patient trials, 99% of 50 patient trials, and 100% of 70 patient trials. Nilotinib trials simulated with 40% between-occasion variability resulted in significant food effects on clearance being retrieved in 50% of 20 patient trials, 78% of 50 patient trials, and 88% of 70 patient trials. The ranges of percent bias and precision of food effect were -2.16% to 11.8% and 6.3% to 38.8%, respectively.
Table 5.4. **Bias and precision of nilotinib oral clearance at the population level and individual level, between-individual variability, and between-occasion variability**

<table>
<thead>
<tr>
<th>Between-occasion variability</th>
<th>Number of patients in trial</th>
<th>Oral Clearance – Population level</th>
<th>Oral Clearance – Individual level</th>
<th>Between-individual variability</th>
<th>Between-occasion variability</th>
<th>Food effect</th>
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<tr>
<td></td>
<td></td>
<td>Bias</td>
<td>Precision</td>
<td>Bias</td>
<td>Precision</td>
<td>Bias</td>
</tr>
<tr>
<td>10%</td>
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<td>-11.8%</td>
<td>16.0%</td>
<td>-0.6%</td>
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<td>50</td>
<td>-12.0%</td>
<td>16.3%</td>
<td>-0.5%</td>
<td>4.3%</td>
<td>-9.9%</td>
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<td>-12.8%</td>
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<td>25%</td>
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<td>40%</td>
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<td>-13.3%</td>
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<td>-0.9%</td>
<td>7.1%</td>
<td>-7.9%</td>
</tr>
</tbody>
</table>
4. Discussion

Virtual clinical trials of abiraterone and nilotinib using sparse concentration measurements and a population PK sampling time design were simulated to test whether the drug exposure of each simulated patient and its variability under different prandial conditions on oral clearance for two recently approved drugs would be retrieved accurately and precisely. The trial design is a typical phase II design in the NCI cooperative system and this assessment provides and evaluation of whether there is value in drawing sparse samples to utilize in population pharmacokinetic analysis for the determination of individual drug exposure. It was important to assess this particular study design as it is widely used and insight on the value of drug concentration sampling under these conditions was unclear. Population as well as individual pharmacokinetic parameters for abiraterone, with a large food effect, and nilotinib, with a smaller food effect, were well estimated from the virtual trial results. This evaluation of whether between-individual and between-occasion variability can be well captured with a significant covariate effect on oral clearance (the underlying food effect) at different levels of variability on anticancer drug exposure is a novel observation. As the prior knowledge of more than 100% between-individual variability was introduced to oral clearance in the abiraterone trial, the model estimation of the population oral clearance parameter and between-individual variability have relatively poor percent bias and precision compared to the nilotinib trial. This is reflected in the power calculation showing that only 10 to 20% of the trials across three between-occasion variability levels (10, 25, 40%) have population oral clearance estimates within 20% of the true value. This finding indicates that retrieving population level effects with large between-individual
variability may need a much larger trial size or more intense sampling schedule or a combination of both.

Between-occasion variability estimates were reasonable considering both percent bias and precision were less than 30% in most trial simulations. However, the percent bias and precision were relatively poor for the simulated abiraterone trials with 20 patients and 10% within-individual variability. One possible explanation of this is that the system is less able to capture between-individual variability if the actual variability is small, but the true mechanism contributing to this poor bias at 10% between-individual variability is as yet unclear.

This model was unable to estimate both residual error and between-occasion variability accurately simultaneously. In order to accurately capture the between-individual variability, the residual error estimation was fixed to published values. This suggests that more than one sample per occasion will be required to distinguish these hierarchies of variability simultaneously.

The effects of two important features of such clinical trials were quantified using a NONMEM based simulation analysis. These were the power to detect differences in oral drug clearance related to the prandial state of the study participants and the degree of between-individual variability and sample size. When the between-individual variability was set to 25% for a trial with 20 patients, 80% of the simulated nilotinib trials resulted in the detection of a statistically significant reduction in oral clearance caused by the food effect. The power to detect this food effect on oral drug clearance increased to 100% when the number of patients per trial was increased from 20 to 70. At 40% between-individual variability, the food effect signal was observed in only 50% of trials with 20
patients, increasing to 88% when the number of patients increased to 50. The percent precision also indicated that the food effect as a covariate was captured less precisely when between-occasion variability increased, and the trial sample size was reduced. In contrast, abiraterone with a much more substantial food effect (92% reduction in oral clearance) resulted in a power to detect the food effect of 100% in the smallest trial evaluated (20 patients). This virtual trial also had 40% between-individual variability and resulted percent bias and precision on the Pop PK parameters that were all less than 20%. Using a modeling approach, we were able to simulate a complex clinical oncology population pharmacokinetic trial setting and capture the between-occasion variability and the magnitude of individual drug exposure in the presence of a large food effect for two recently approved oral anti-cancer agents. This simulated approach facilitated an early evaluation of the proposed trial design. However, clinical trial simulations are generated based on many trial assumptions, and these assumptions may include uncertainty. In fact, with different underlying assumptions, the simulated outcomes can differ, so multiple scenarios as well as assumptions must be tested in order to fully interpret the relevance of the results.

There are limitations to this analysis. First, a dropout model was not incorporated in this study design which can potentially contribute to censor events in a 3 month study. The simulations also assumed parameter distributions based on available food effect assessment in previous non-compartmental analysis. Because of the established sampling schedule, both drugs were better estimated with a simplified model structure. Despite this, the model provides an approximation of the actual behavior and can capture the trend of the variability within the population. In fact, this simplified modeling approach
was previously proposed to assess the pharmacokinetics of some drug entities in the trials with a relative sparse sampling schedule (Scerwin et al., 2012; van Erp et al., 2011). The sampling schedule could also be optimized for the identification of food effects and model parameters, but the objective of this work was to evaluate trial designs given a commonly used sampling schedule. Second, non-compliance was not considered in this study design as we assumed that the compliance rate should be reasonably controlled in the clinical trial although this will result in a higher residual error and between-individual variability than compliance accounted for using electronically monitoring (Vrijens et al., 2005). This study emphasizes the importance of addressing trial designs where intensive sampling cannot be obtained and yet there is a need to understand the drug exposure characteristics to what are otherwise medications with highly variable pharmacokinetic disposition.
CHAPTER VI: Genetic determinants of vincristine neuropathy in pediatric acute lymphoblastic leukemia patients

1. Introduction

Vincristine is one of the most commonly prescribed anticancer agents, and it has been used to treat a wide variety of malignancies. Even though the drug is commonly used, the dose optimization of vincristine and potential causes of severe side effects and lack of efficacy still remain unclear. In fact, vincristine plays a role in many cellular mechanisms. It not only binds irreversibly to microtubules and spindle proteins and interferes with the formation of the mitotic spindle, but also depolymerizes microtubules and may also interfere with amino acid, cyclic AMP, and glutathione metabolism. In addition, it is associated with calmodulin-dependent Ca++-transport ATPase activity, cellular respiration, and nucleic acid and lipid biosynthesis (National Cancer Institute, 2011).

Similar to other chemotherapy agents such as taxanes and platinums, vincristine is associated with highly variable cumulative dose-dependent neurotoxicity, secondary to its binding to tubulin dimers, which can inhibit the assembly of microtubule and arrest mitosis in metaphase (Park et al., 2013; Howlader et al., 2012; Verstappen et al., 2005; Postma et al., 1993; Haim et al., 1994). In severe cases, neurotoxicity may lead to dose reduction, which may negatively impact the drug efficacy. In addition, substantial variability in vincristine pharmacokinetics with up to a 40-fold interpatient variation has been reported in the literature (Crom et al., 1994; de Graaf et al., 1995), and this pharmacokinetic variability may be associated with severity of neurotoxicity. Published data also indicate that vincristine pharmacokinetics may in fact be associated with long-
term outcomes in children of acute lymphoblastic leukemia with rapid clearance being associated with a greater risk of relapse. (Lonnerholm et al., 2008).

The variability of vincristine pharmacokinetics might be potentially explained in part by genetic differences in cytochrome P450 3A family enzymes, which are important for vincristine metabolism (Dennison, et al., 2006).

As previous study shown, Caucasian children with ALL more commonly suffer from vincristine-induced peripheral neuropathy (VIPN) than do children of other ethnicities due to CYP3A5 genotype (Dennison et al., 2006). In fact, another study suggests that polymorphisms of multi-drug resistance (MDR1) gene which code as an efflux transporter are associated with peripheral neuropathy (Te Loo et al., 2010). This might also contribute to the variability of vincristine PK, and the genetic impact on drug toxicity. In addition, study published by Hertz et al. suggested that CYP2C8 polymorphisms can increase the risk of neuropathy in breast cancer patients treated with paclitaxel (Hertz et al., 2013). This indicated that in the case of paclitaxel drug toxicity may be associated with pharmacokinetic exposure through metabolic pathways. However, those findings are controversial across studies with limited validation in the literature (Hertz et al., 2014). In a study of 329 subjects by Broyl et al., an early-onset vincristine-induced neuropathy was characterized by the up-regulation of genes involved in cell cycle and proliferation, including AURKA and MKI67, and also by the presence of single-nucleotide polymorphisms (SNPs) in genes involved in these processes, such as GLI1 (rs2228224 and rs2242578) (Broyl et al., 2010). Similar to the vincristine association analyses, many studies have aimed to identify potential genetic risks in predicting neuropathy induced by platinum agents or Taxanes. In those genomic studies,
few candidate genes (FGD4, TUBB2A, ARHGEF and PRX) were found to be associated with paclitaxel-induced peripheral neurotoxicity. These results indicated that drug-induced peripheral neurotoxicity may share genetic roots as hereditary neuropathies (Baldwin et al., 2012; Leandro-Garcia et al., 2012; Beutler et al., 2014; Travis et al., 2014). Furthermore, McWhinney-Glass’s group further explored genetic risk of four SNPs (SOX10, BCL2, OPRM1 and TRPV1) in an association analysis of platinum/taxane-induced neurotoxicity. Interestingly, a cumulative association of these genetic variations with neurotoxicity was observed in this population of 404 patients with ovarian disease (McWhinney-Glass et al., 2013).

In order to better understand VIPN and predict risk for toxicity to ultimately aid in vincristine dose optimization and to further explore the findings from the previous studies, a genome-wide association analysis was proposed to take an unbiased approach in evaluating the association between germline variants and VIPN in pediatric ALL patients. Indeed, as the genomics technologies improve, genetic biomarkers provide a unique tool for prediction of disease outcomes of interest. The objective of this study is to build a candidate gene list including polymorphisms associated with VIPN from the GWAS study as well as other association analysis approaches. The rationale for the proposed research is that once we understand the impact of specific pharmacogenetic changes on vincristine toxicity, we will be able to use this information to optimize dosing of this important drug for individual patients.
2. Materials and Methods

2.1 ALL Patient Population and Study Design

A total of 2154 children with precursor B cell acute lymphoblastic leukemia from the Pediatric Oncology Group (POG) 9904, 9905 and 9906 trials were enrolled. In conjunction with 9905, the objective of the 9904 trial was to determine if a delayed multi-drug intensification, administered in the context of intensive anti-metabolite therapy improved outcomes and compare the different durations of methotrexate infusion. 9906 was aimed at specifically evaluating patients at high risk for treatment failure. Those who had Down’s syndrome, Charcot Marie Tooth disease, baseline peripheral neuropathy, or a history of liver disease with chronic elevation in liver function tests to greater than 5-times the upper limit of normal based on normal values for age were excluded from this GWAS. Genotyping of germline samples from 1888 patients on the 9900 trials has been completed using the Affymetrix GeneChip Human Mapping 6 and 500K arrays for 9904/9905 and 9906, respectively. The number of vincristine doses per patient over the course of treatment ranged from 18-23, depending on specific protocol and treatment arm. Standard vincristine dosing of 1.5mg/m²/dose was used. The complete treatment duration is 2.5 years from the date of diagnosis for females and 3.5 years for males. The treatment plan is summarized in Figure 6.1.
**Vincristine dose regimen scheme in POG 9904 trial**

| Day | 0 | 7 | 14 | 21 | 28 | 35 | 42 | 49 | 56 | 63 | 70 | 77 | 84 | 91 | 98 | 105 | 112 | 119 | 126 | 133 | 140 | 147 | 154 | 161 | 168 | 175 | 182 | 189 | 196 | 203 | 210 | 217 | 224 | 231 | 238 |
|-----|---|---|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|
| Week 1 | 0 | 7 | 14 | 21 | 28 | 35 | 42 | 49 | 56 | 63 | 70 | 77 | 84 | 91 | 98 | 105 | 112 | 119 | 126 | 133 | 140 | 147 | 154 | 161 | 168 | 175 | 182 | 189 | 196 | 203 | 210 | 217 | 224 | 231 | 238 |

**Vincristine dose regimen scheme in POG 9905 trial**

| Day | 0 | 7 | 14 | 21 | 28 | 35 | 42 | 49 | 56 | 63 | 70 | 77 | 84 | 91 | 98 | 105 | 112 | 119 | 126 | 133 | 140 | 147 | 154 | 161 | 168 | 175 | 182 | 189 | 196 | 203 | 210 | 217 | 224 | 231 | 238 |
|-----|---|---|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|
| Week 1 | 0 | 7 | 14 | 21 | 28 | 35 | 42 | 49 | 56 | 63 | 70 | 77 | 84 | 91 | 98 | 105 | 112 | 119 | 126 | 133 | 140 | 147 | 154 | 161 | 168 | 175 | 182 | 189 | 196 | 203 | 210 | 217 | 224 | 231 | 238 |
Vincristine dose regimen scheme in POG 9906 trial

Figure 6.1. Vincristine dosage regimens for POG 9904, 9905 and 9906 trials.
2.2 Genotype Data and Quality Control

DNA(500ng) was digested with restriction enzymes, amplified, labeled, and hybridized to the Affymetric GeneChip Human Mapping 6 set for P9004 and 9005 and 500K for 9006. The genotypic data included raw data files, genotypes, quality scores, intensity values, SNPs and sample summary tables. QC was performed to remove both samples and markers which were unreliable in the following fashion. Samples with >5% missing rate were excluded. Furthermore, SNPs with a study-wide missing data rate of >5% and/or evidence of Hardy-Weinberg disequilibrium (p≤ 0.0001) were discarded. SNPs with minor allele frequency (MAF) of <0.05 were also removed from the analysis because previous studies have shown that these SNPs have little power to detect association and are more prone to genotypic errors resulting in false positive evidence of association. In addition, MAF < 0.10 cutoff was applied for the recessive model in the analysis. PLINK was proposed to manipulate the data by generating both per sample and per SNP metrics to assess the quality of the genotypic data (Purcell et al., 2007).

2.3 Population Stratification

A principal components approach (Price et al., 2006) was applied to correct for any possible stratification errors. A subset of the genome-wide SNP data was used to identify components that reflect population structure. Scores for each individual from the initial principal component assessment were calculated and included as covariates in the subsequent association analyses.

2.4 Phenotype Identification

Vincristine is associated with highly variable cumulative dose-dependent peripheral neurotoxicity that often necessitates chemotherapy dose reductions, thereby
compromising efficacy. To do this association analysis, the first step was to define our phenotype of interest. National Cancer Institute Common Terminology Criteria for Adverse Events, version 3.0 (CTCAE v3.0) was used to capture and grade adverse events in this study. We defined neuropathy for the purposes of this analysis as any sensory or motor neuropathy ≥ grade 2. Neuropathic pain was also captured in this study. For the purpose of this analysis, we designate neuropathic pain as secondary neuropathy as any sensory neuropathy, motor neuropathy or neuropathic pain ≥ grade 2.

2.5 Genome-Wide Association Study

The analysis started with association analysis of genotype frequencies with primary neuropathy (CTCAT V3.0 sensory or motor neuropathy). Neuropathy was treated as a binary variable; ie, 1 for having neuropathy and 0 for no neuropathy. Clinical, demographic, and population stratification variables were tested as covariates in the regression analyses. Genotype frequencies were tested in dominant, recessive and additive (gene-dose) model. This strategy was also used to test the association between SNPs and secondary neuropathy. These analyses were performed in R coxph package (Therneau et al., 2014). The corresponding p-value of each SNP was summarized across chromosomes in Manhattan and QQ plots.

2.6 Gene Enrichment Analysis

The gene enrichment analysis was done by focusing on enrichment of specific target genes (Shamir, 2010). In this case, the target genes are genes previously identified as being related to neuropathy or pain. The initial step of the analysis was to obtain SNP rsIDs along with their genomic positions and p-value from the SNP6.0 Affymetrix COG data which was previously analyzed in the survival analysis with neuropathy phenotype.
SNPs were annotated with their respective genes using their positions and with the annotation file from UCSC genome browser. This was done by matching the SNP chromosome number with the gene chromosome number and then looking for SNPs that within the gene base pair co-ordinates. Upstream or downstream regions of the genes were not considered. Using the significance frequency table, a Fisher’s exact test was run to calculate the odds ratio based on equation 1 for each gene.

<table>
<thead>
<tr>
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<th>P-value &lt;0.05</th>
<th>P-value &gt;0.05</th>
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<td>Candidate gene</td>
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<td>N12</td>
</tr>
<tr>
<td>Other genes</td>
<td>N21</td>
<td>N22</td>
</tr>
</tbody>
</table>

\[
\text{Odds Ratio} = \frac{N11 \times N22}{N12 \times N21} \quad \text{(Equation 1)}
\]

N is the number of SNPs in each category. The p-value was estimated in Fisher’s extract test. Based on the criteria of p-value <0.05 from Fisher’s extract test and calculated odds ratio >1, genes which passed the criteria were selected.

### 2.7 Pathway Analysis

Significant pathways in biological processes were identified in an interactive pathway analysis from Ingenuity systems (Ingenuity systems analysis). The following steps were used in the analysis: “(1) Genes identified as significant from the experimental data sets were overlaid onto the interactome. Focus genes re-identified as the subset having direct interaction(s) with other genes in the database. (2) The specificity of connections for each focus gene was calculated by the percentage of its connections to other significant genes. The initiation and growth of pathways proceeded from genes with the highest specificity of connections. Each pathway had a maximum of 35 genes. (3)
Pathways of highly interconnected genes were identified by statistical likelihood using the following equation:

\[
\text{Score} = -\log_{10}(1 - \sum_{i=0}^{f-1} \frac{C(G,i)C(N-G,S-i)}{C(N,s)})
\]

where \( N \) is the number of genes in the genomic network, of which \( G \) are focus genes, for a pathway of genes, \( f \) of which are focus genes, and \( C(n,k) \) is the binomial coefficient. (4) Pathways with a score greater than 4 (\( P < 0.0001 \)) were combined to form a composite network representing the underlying biology of the process.”

2.8 Gene Expression Analysis

Gene expression omnibus (GEO) is a public functional genomics data repository supporting MIAME-compliant data submissions (Lehnhardt et al., 2005). To explore whether the gene expression level was changed after treating with vincristine, a keyword search of vincristine in GEO database was performed. SNPs in COG data were annotated with their respective genes using their positions and with the annotation file from UCSC genome browser. These genes were compared with the genes in GEO. Those overlapped genes were further filtered by the gene expression level with a threshold of greater than 50% up-regulation or down-regulation of the vincristine treated replicates compared to the control. For those genes which passed the threshold, the SNPs with their p-values from the GWAS study were evaluated and visualized in the QQ plot.

2.9 eQTL (GTEx) Analysis

In addition to the GEO database, tissue data from GTEx (Genotype-Tissue Expression) project (Gibbs et al., 2010) was explored to determine the correlations between genotype and the tissue-specific gene expression level. This could help to identify the expression quantitative trait locus (eQTL) that might regulate the expression level of mRNAs or proteins. Based on the available data from the published tissue bank,
four of the brain tissues including cerebellum, frontal cortex, temporal cortex and pons were considered in the correlation analysis with neuropathy.

3. Results

3.1 ALL Patient Characteristics

Within 2154 ALL patients enrolled in POG 9904, 9905 and 9906 trials, genotyping of 1888 individuals were eligible. To further insure the genotyping quality, only patients from POG 9904 and 9905 who were genotyped using Affymetrix 6.0 were included in the initial analysis. Furthermore, a principle component approach was applied to correct the population stratification error by using Hapmap references of Han Chinese in Beijing, China (CHB), Utah residents with Northern and Western European ancestry from the CEPH collection (CEU) and African ancestry in Southwest USA (ASW). Based on the result of the two principal component analyses (PC1 and PC2) in Figure 6.2, Caucasians were better overlapped with the reference and represented the majority of the population. In addition, data which did not pass the additional quality control including samples or SNP with >5% missing data rate and/or evidence of Hardy-Weinberg disequilibrium (p≤ 0.0001) and/or MAF <0.05 for dominant and additive model and/or MAF< 0.10 for recessive model were excluded, and a total of 587,014 SNPs of 1068 individuals remained in the analysis (Figure 6.3).
Figure 6.2. Flowchart of ALL patient selection in the final GWAS analysis.  
MAF: minor allele frequency; dom: dominant model; add: additive model; rec: recessive model;  
HW: Hardy Weinberg equilibrium.
Figure 6.3. **Population stratification using principal component analysis.** HapMap reference was used to compare the data. CHB: Han Chinese in Beijing, China. CEU: Utah residents with Northern and Western European ancestry from the CEPH. ASW: African ancestry in Southwest USA.
3.2 Identification of Genomic Loci Associated with ALL Neuropathy Risk in GWAS Analysis

The initial step of the analysis coded homozygous major alleles, heterozygous, and homozygous minor alleles as 0, 1 and 2, respectively using PLINK. After the data was manipulated, a Cox proportional hazard model was performed to test association between markers and the primary and the secondary neuropathy in three genetic models. The p-values in –log10 were shown across chromosomes in Manhattan plot and QQ plot for primary neuropathy and secondary neuropathy in Figure 6.4 A, B, C, D, E and F and Figure 6.5A, B, C, D, E and F, respectively.
Figure 6.4. Manhattan plot and QQ plot of GWAS association analysis with primary neuropathy.

A and B: dominant model; C and D: recessive model; E and F: additive model.
Figure 6.5. Manhattan plot and QQ plot of GWAS association analysis with secondary neuropathy.
A and B: dominant model; C and D: recessive model; E and F: additive model.
In order to determine the candidate SNPs for validation, a threshold of p-value of $1 \times 10^{-4}$ was selected. Those candidate SNPs with p-value $< 1 \times 10^{-5}$ were further investigated and identified the location of gene region and visualized closely in zoom-in plot (Figure 6.6). A total of 23 genes were identified across both neuropathy phenotypes and three genetic model structures. The rsID, position, minor allele, MAF and hazard ratio of those candidate SNPs as well as its corresponding gene were summarized in Table 6.1.
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<th>Zoom-in plot</th>
<th>Corresponding SNPs</th>
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</tr>
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<td><strong>SNP</strong></td>
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Figure 6.6. Zoom-in plots of each candidate gene from GWAS study and their corresponding SNPs. NE: neuropathy (primary neuropathy); NEP: neuropathy and neuropathic
pain (secondary neuropathy); DOM: dominant model; REC: recessive model; DOSE: additive model.
Table 6.1. Candidate genes selected from GWAS study.

<table>
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<tr>
<th>SNP</th>
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<th>Position</th>
<th>Gene</th>
<th>Genetic model</th>
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### Top SNPs from GWAS to vincristine induced secondary neuropathy

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<th>Position</th>
<th>Gene</th>
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<th>Minor Allele</th>
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<td>Additive</td>
<td>C</td>
<td>0.19</td>
<td>1.947</td>
<td>2.37E-06</td>
</tr>
<tr>
<td>rs290505</td>
<td>5</td>
<td>60307964</td>
<td>NDUFAF2</td>
<td>Additive</td>
<td>T</td>
<td>0.20</td>
<td>0.532</td>
<td>2.53E-06</td>
</tr>
<tr>
<td>rs162231</td>
<td>5</td>
<td>60296556</td>
<td>NDUFAF2</td>
<td>Additive</td>
<td>T</td>
<td>0.20</td>
<td>1.887</td>
<td>4.01E-06</td>
</tr>
<tr>
<td>rs158816</td>
<td>5</td>
<td>60245145</td>
<td>NDUFAF2</td>
<td>Additive</td>
<td>G</td>
<td>0.20</td>
<td>0.545</td>
<td>9.36E-06</td>
</tr>
<tr>
<td>rs158563</td>
<td>5</td>
<td>60255004</td>
<td>NDUFAF2</td>
<td>Additive</td>
<td>T</td>
<td>0.34</td>
<td>1.796</td>
<td>1.38E-05</td>
</tr>
<tr>
<td>rs292614</td>
<td>5</td>
<td>60304171</td>
<td>NDUFAF2</td>
<td>Additive</td>
<td>T</td>
<td>0.17</td>
<td>1.802</td>
<td>1.47E-05</td>
</tr>
<tr>
<td>rs167912</td>
<td>5</td>
<td>60284996</td>
<td>NDUFAF2</td>
<td>Additive</td>
<td>T</td>
<td>0.20</td>
<td>0.558</td>
<td>1.64E-06</td>
</tr>
<tr>
<td>rs162240</td>
<td>5</td>
<td>60299344</td>
<td>NDUFAF2</td>
<td>Additive</td>
<td>G</td>
<td>0.35</td>
<td>1.796</td>
<td>1.97E-05</td>
</tr>
<tr>
<td>rs11951504</td>
<td>5</td>
<td>60372438</td>
<td>NDUFAF2</td>
<td>Additive</td>
<td>G</td>
<td>0.17</td>
<td>1.745</td>
<td>5.17E-05</td>
</tr>
<tr>
<td>rs4647108</td>
<td>5</td>
<td>60197715</td>
<td>ERCC8</td>
<td>Additive</td>
<td>G</td>
<td>0.21</td>
<td>1.849</td>
<td>5.62E-06</td>
</tr>
<tr>
<td>rs158928</td>
<td>5</td>
<td>60227923</td>
<td>ERCC8</td>
<td>Additive</td>
<td>T</td>
<td>0.21</td>
<td>1.837</td>
<td>7.64E-06</td>
</tr>
<tr>
<td>rs4647078</td>
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<td>60217038</td>
<td>ERCC8</td>
<td>Additive</td>
<td>G</td>
<td>0.20</td>
<td>1.788</td>
<td>2.02E-05</td>
</tr>
<tr>
<td>rs4700398</td>
<td>5</td>
<td>60135985</td>
<td>ELOVL7</td>
<td>Additive</td>
<td>G</td>
<td>0.26</td>
<td>1.723</td>
<td>2.34E-05</td>
</tr>
<tr>
<td>rs6869332</td>
<td>5</td>
<td>60129361</td>
<td>ELOVL7</td>
<td>Additive</td>
<td>T</td>
<td>0.20</td>
<td>0.567</td>
<td>2.74E-05</td>
</tr>
<tr>
<td>rs6877849</td>
<td>5</td>
<td>60088619</td>
<td>ELOVL7</td>
<td>Additive</td>
<td>G</td>
<td>0.21</td>
<td>0.582</td>
<td>7.5E-05</td>
</tr>
<tr>
<td>rs6449502</td>
<td>5</td>
<td>60093095</td>
<td>ELOVL7</td>
<td>Additive</td>
<td>T</td>
<td>0.21</td>
<td>0.581</td>
<td>8.42E-05</td>
</tr>
<tr>
<td>rs5017492</td>
<td>3</td>
<td>17544747</td>
<td>NAALADL2</td>
<td>Recessive</td>
<td>T</td>
<td>0.27</td>
<td>3.176</td>
<td>3.12E-06</td>
</tr>
<tr>
<td>rs6792776</td>
<td>3</td>
<td>17549067</td>
<td>NAALADL2</td>
<td>Recessive</td>
<td>G</td>
<td>0.25</td>
<td>2.944</td>
<td>1.29E-05</td>
</tr>
<tr>
<td>rs3768125</td>
<td>1</td>
<td>23690181</td>
<td>ACTN2</td>
<td>Dominant</td>
<td>G</td>
<td>0.42</td>
<td>0.502</td>
<td>7.48E-05</td>
</tr>
<tr>
<td>rs17091414</td>
<td>10</td>
<td>115962953</td>
<td>TDRD1</td>
<td>Additive</td>
<td>G</td>
<td>0.23</td>
<td>1.801</td>
<td>2.33E-05</td>
</tr>
<tr>
<td>rs2616657</td>
<td>10</td>
<td>115969622</td>
<td>TDRD1</td>
<td>Additive</td>
<td>G</td>
<td>0.17</td>
<td>1.724</td>
<td>7.01E-05</td>
</tr>
<tr>
<td>rs17144752</td>
<td>7</td>
<td>122359692</td>
<td>CDPS2</td>
<td>Recessive</td>
<td>G</td>
<td>0.12</td>
<td>1.38</td>
<td>1.29E-05</td>
</tr>
<tr>
<td>rs7777183</td>
<td>7</td>
<td>122316632</td>
<td>CDPS2</td>
<td>Recessive</td>
<td>C</td>
<td>0.30</td>
<td>2.897</td>
<td>4.19E-05</td>
</tr>
<tr>
<td>rs1522271</td>
<td>3</td>
<td>191996779</td>
<td>FGF12</td>
<td>Recessive</td>
<td>G</td>
<td>0.43</td>
<td>0.865</td>
<td>1.39E-05</td>
</tr>
<tr>
<td>rs6793796</td>
<td>3</td>
<td>191977079</td>
<td>FGF12</td>
<td>Recessive</td>
<td>G</td>
<td>0.41</td>
<td>0.831</td>
<td>7.08E-05</td>
</tr>
<tr>
<td>rs10203853</td>
<td>2</td>
<td>234687418</td>
<td>MROH2A</td>
<td>Additive</td>
<td>T</td>
<td>0.24</td>
<td>0.943</td>
<td>2.23E-06</td>
</tr>
<tr>
<td>rs10209214</td>
<td>2</td>
<td>234687583</td>
<td>MROH2A</td>
<td>Additive</td>
<td>C</td>
<td>0.34</td>
<td>0.606</td>
<td>5.07E-05</td>
</tr>
</tbody>
</table>
3.3 Identification of Genomic Loci Associated with ALL Neuropathy Risk in Gene-Enrichment Analysis

A gene-enrichment analysis based on a bottom-up approach was conducted to explore genes previously identified as drug targets for treating neuropathy or pain. A total of 101 genes were identified for evaluation in this analysis. Among these 101 target genes, the SNPs were enriched in the gene regions were tested using Fisher’s extract test. The threshold of significance was set to be p-value at 0.05. A total of eight genes including CACNA1D, SLC29A4, CACNA1C, GRIK1, SCN8A, CACNB1, GRIN3A and SLC22A1 with odds ratio greater than 1 were identified and listed in Table 6.2. The zoom-in plots of those genes with their p-values were summarized in Figure 6.7.
<table>
<thead>
<tr>
<th>Gene</th>
<th>Length (bp)</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>CACNA1D</td>
<td>317,810</td>
<td>53,528,683-53,846,492</td>
</tr>
<tr>
<td>SLC29A4</td>
<td>29,705</td>
<td>5,314,000-5,343,704</td>
</tr>
<tr>
<td>CACNA1C</td>
<td>727,164</td>
<td>2,079,952-2,807,115</td>
</tr>
<tr>
<td>GRIK1</td>
<td>403,029</td>
<td>30,909,254-31,312,282</td>
</tr>
<tr>
<td>SCN8A</td>
<td>218,258</td>
<td>51,984,050-52,202,307</td>
</tr>
<tr>
<td>CACNB1</td>
<td>24,248</td>
<td>37,329,709-37,353,956</td>
</tr>
<tr>
<td>GRIN3A</td>
<td>169,229</td>
<td>104,331,634-104,500,862</td>
</tr>
<tr>
<td>SLC22A1</td>
<td>36,930</td>
<td>160,542,821-160,579,750</td>
</tr>
</tbody>
</table>

Table 6.2. **Candidate genes selected from gene-enrichment analysis.**
<table>
<thead>
<tr>
<th>Gene</th>
<th>P-value</th>
<th>Odds ratio</th>
<th>Zoom-in plot</th>
</tr>
</thead>
<tbody>
<tr>
<td>CACNAID</td>
<td>2.96E-10</td>
<td>4.405549</td>
<td><img src="image1" alt="Plot" /></td>
</tr>
<tr>
<td>SLC29A4</td>
<td>0.000133</td>
<td>24.02168</td>
<td><img src="image2" alt="Plot" /></td>
</tr>
<tr>
<td>CACNA1C</td>
<td>0.004477</td>
<td>1.894156</td>
<td><img src="image3" alt="Plot" /></td>
</tr>
<tr>
<td>GRIK1</td>
<td>0.008791</td>
<td>2.109875</td>
<td><img src="image4" alt="Plot" /></td>
</tr>
<tr>
<td>Gene</td>
<td>P-value</td>
<td>Odds Ratio</td>
<td></td>
</tr>
<tr>
<td>---------</td>
<td>----------</td>
<td>------------</td>
<td></td>
</tr>
<tr>
<td>SCN8A</td>
<td>0.016998</td>
<td>2.746166</td>
<td></td>
</tr>
<tr>
<td>CACNB1</td>
<td>0.024969</td>
<td>19.22312</td>
<td></td>
</tr>
<tr>
<td>GRIN3A</td>
<td>0.038966</td>
<td>2.089492</td>
<td></td>
</tr>
<tr>
<td>SLC22A1</td>
<td>0.045554</td>
<td>4.805872</td>
<td></td>
</tr>
</tbody>
</table>

Figure 6.7. **Zoom-in plots of each candidate gene from Gene enrichment analysis study and their corresponding P-value and odds ratio.**
3.4 Pathway Identification of Genomic Loci Associated with ALL Neuropathy Risk

The candidate genes selected from the association analysis were further evaluated in an interactive pathway analysis. Only one pathway, pregenolone biosynthesis, passed the p-value threshold. In addition, two networks were selected based on the score that the system calculated. The first network had a score of 35 with 14 focus moles including ACTN2, CNTNAP2, CSNK1G, ERCC8, ERLEC1, FGF12, GRM8, MICAL3, NAALADL2, NDUFAF2, NEK6, NOD1, PHACTR1 and TDRD1; it is cell-to cell signaling and interaction, infectious disease, inflammatory (Figure 6.8). The second network with a score of 15 with 7 focus moles including CADPS2, ELOVL7, FHIT, LPPR5, PDE8B, PRKG1 and ZFPM2 is tissue development, tumor morphology, cancer (Figure 6.9).
Figure 6.8. The network of cell-to-cell signaling and interaction, infectious disease and inflammatory. 14 genes included ACTN2, CNTNAP2, CSNK1G, ERCC8, ERLEC1, FGF12, GRM8, MICAL3, NAALADL2, NDUFAF2, NEK6, NOD1, PHACTR1 and TDRD1 were identified in this network.
Figure 6.9. The network of tissue development, tumor morphology and cancer.

7 genes including CADPS2, ELOVL7, FHT, LPPR5, PDE8B, PRKG1 and ZFPM2 were identified in this network.
3.5 The Change in Gene Expression of Genomic Loci Associated with ALL Neuropathy Risk

The correlation of genetic information and toxicity was further evaluated at the gene expression level to consider whether the genes of interest are functional and expressed. A study of a fibrosarcoma cell line response to various cytostatic drugs was utilized in which the control cell line was compared to that (Figure 6.10). The gene expression level of the sample treated with vincristine was normalized by the control. A total of 4619 and 16893 SNPs reached the threshold of greater than 50% down-regulation and up-regulation, respectively. Based on the QQ plots, most of the p-values of SNPs were shown to approximately lie on the line $y = x$ meaning that the p-values of the SNPs are likely to be randomly distributed. Few SNPs were deviated from the line $y=x$. These are summarized in Figures 6.11 and 6.12 with their corresponding genes.
Figure 6.10. Fibrosarcoma cell line response to various cytostatic drugs in the GEO database. The cell line was treated with Vincristine, doxorubicin or actinomycin.
<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Dominant</th>
<th>Additive</th>
<th>Recessive</th>
<th>Expression level</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary neuropathy</td>
<td><img src="image1" alt="Graph" /></td>
<td><img src="image2" alt="Graph" /></td>
<td><img src="image3" alt="Graph" /></td>
<td>50% Under-Expression</td>
</tr>
<tr>
<td>Primary neuropathy</td>
<td><img src="image4" alt="Graph" /></td>
<td><img src="image5" alt="Graph" /></td>
<td><img src="image6" alt="Graph" /></td>
<td>50% Over-expression</td>
</tr>
<tr>
<td>Secondary neuropathy</td>
<td><img src="image7" alt="Graph" /></td>
<td><img src="image8" alt="Graph" /></td>
<td><img src="image9" alt="Graph" /></td>
<td>50% Under-Expression</td>
</tr>
<tr>
<td>Secondary Neuropathy</td>
<td><img src="image10" alt="Graph" /></td>
<td><img src="image11" alt="Graph" /></td>
<td><img src="image12" alt="Graph" /></td>
<td>50% Over-expression</td>
</tr>
</tbody>
</table>

Figure 6.11. *QQ-plots of genomic loci associated with ALL neuropathy risk in fibrosarcoma cell line.*
<table>
<thead>
<tr>
<th>Phenotype</th>
<th>Expression level</th>
<th>Genetic model</th>
<th>QQ-plot</th>
<th>SNP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary neuropathy</td>
<td>50% Under-Expression</td>
<td>Dominant</td>
<td><img src="image" alt="QQ-plot" /></td>
<td>rs2238034 CACNA1C, rs6882294 ATG10, rs7696226 OTUD4, rs7712216 ATG10</td>
</tr>
<tr>
<td>Primary neuropathy</td>
<td>50% Over-Expression</td>
<td>Dominant</td>
<td><img src="image" alt="QQ-plot" /></td>
<td>rs1867353 MICAL3, rs2075455 MICAL3, rs2165971 MICAL3, rs2167143 KCNA81</td>
</tr>
<tr>
<td>Secondary neuropathy</td>
<td>50% Under-Expression</td>
<td>Recessive</td>
<td><img src="image" alt="QQ-plot" /></td>
<td>rs4543885 NEBL, rs6751734 DNAH7, rs8119756 ADA</td>
</tr>
<tr>
<td>Secondary neuropathy</td>
<td>50% Over-expression</td>
<td>Additive</td>
<td><img src="image" alt="QQ-plot" /></td>
<td>rs4647078 ERCC8, rs4647108 ERCC8</td>
</tr>
<tr>
<td>Secondary neuropathy</td>
<td>50% Over-expression</td>
<td>Recessive</td>
<td><img src="image" alt="QQ-plot" /></td>
<td>rs2717425 PTPRB</td>
</tr>
</tbody>
</table>

Figure 6.12. *Genes and corresponding SNPs of genomic loci associated with ALL neuropathy risk in fibrosarcoma cell line*
3.6 Genomic Loci Associated with ALL Neuropathy Risk in Genotype-Tissue Expression eQTL

In addition to the GEO database, the correlation was also tested in the GTEx database. In this database, mRNA expressions were estimated in the different tissues. Based on current available tissues in the database, this study was carried out to analyze data from four brain tissues which are potentially related to our phenotype of interest. After matching the SNPs between our data and the GTEx database, p-values of these matched SNPs were further visualized in QQ plots across three genetic models and two phenotypes in Figure 6.13. Similar to GEO strategy, the SNPs with deviated –log10 p-values from the line $y = x$ were selected. eQTLs can be described as being cis, where the genotyped marker is within 2 MB of the expressed gene, or trans, in which the genotyped marker is far away from the expressed gene or even on another chromosome (Gibbs et al., 2010). The corresponding genes of the selected SNPs were based on the probe position where the gene expression level was measured in the original eQTL study. As Figure 6.14 showed, in the brain cerebellum tissue, rs10153783 was found to correlate to the gene expression of collectin sub-family member 11 (COLEC11) in the dominant model for both primary and secondary neuropathies. Another SNP, rs359436, is associated with hydroxysteroid (17-beta) dehydrogenase 4 (HSD17B4) gene expression in both dominant and additive models for the secondary neuropathy.
<table>
<thead>
<tr>
<th>Tissue</th>
<th>Phenotype</th>
<th>Dominant</th>
<th>Additive</th>
<th>Recessive</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brain Cerebellum</td>
<td>Primary neuropathy</td>
<td><img src="image1" alt="Graph" /></td>
<td><img src="image2" alt="Graph" /></td>
<td><img src="image3" alt="Graph" /></td>
</tr>
<tr>
<td>Brain Cerebellum</td>
<td>Secondary neuropathy</td>
<td><img src="image1" alt="Graph" /></td>
<td><img src="image2" alt="Graph" /></td>
<td><img src="image3" alt="Graph" /></td>
</tr>
<tr>
<td>Brain Front Cortex</td>
<td>Primary neuropathy</td>
<td><img src="image1" alt="Graph" /></td>
<td><img src="image2" alt="Graph" /></td>
<td><img src="image3" alt="Graph" /></td>
</tr>
<tr>
<td>Brain Front Cortex</td>
<td>Secondary neuropathy</td>
<td><img src="image1" alt="Graph" /></td>
<td><img src="image2" alt="Graph" /></td>
<td><img src="image3" alt="Graph" /></td>
</tr>
<tr>
<td>Brain Temporal Cortex</td>
<td>Primary neuropathy</td>
<td><img src="image1" alt="Graph" /></td>
<td><img src="image2" alt="Graph" /></td>
<td><img src="image3" alt="Graph" /></td>
</tr>
<tr>
<td>Brain Temporal Cortex</td>
<td>Secondary neuropathy</td>
<td><img src="image1" alt="Graph" /></td>
<td><img src="image2" alt="Graph" /></td>
<td><img src="image3" alt="Graph" /></td>
</tr>
</tbody>
</table>
Brain Pons  Primary neuropathy

Brain Pons  Secondary neuropathy

Figure 6.13. QQ-plots of genomic loci associated with ALL neuropathy risk in eQTL data of four brain tissues.
<table>
<thead>
<tr>
<th>Tissue</th>
<th>Phenotype</th>
<th>Genetic model</th>
<th>QQ-plot</th>
<th>eQTL probe gene</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brain Cerebellum</td>
<td>Primary neuropathy</td>
<td>Dominant</td>
<td><img src="image1.png" alt="QQ-plot" /></td>
<td>COLEC11</td>
</tr>
<tr>
<td>Brain Cerebellum</td>
<td>Secondary neuropathy</td>
<td>Dominant</td>
<td><img src="image2.png" alt="QQ-plot" /></td>
<td>COLEC11</td>
</tr>
<tr>
<td>Brain Frontal Cortex</td>
<td>Secondary neuropathy</td>
<td>Dominant</td>
<td><img src="image3.png" alt="QQ-plot" /></td>
<td>HSD17B4</td>
</tr>
<tr>
<td>Brain Frontal Cortex</td>
<td>Secondary neuropathy</td>
<td>Additive</td>
<td><img src="image4.png" alt="QQ-plot" /></td>
<td>HSD17B4</td>
</tr>
</tbody>
</table>

Figure 6.14. *Indirect association of ALL neuropathy risk with eQTL expression.*
4. Discussion

Based on a previous study from Dennison and colleagues, they concluded that CYP3A5 expressers experienced less vincristine-induced peripheral neuropathy and had lower metabolic ratios compared to CYP3A5 non-expressers (Dennison, et al., 2006). This finding stimulates our interests in identifying potential genetic explanations for the difference in the pharmacokinetic exposures in a large population and identifying associations between genetic markers and drug-induced neuropathy. In fact, a subgroup of patients had to discontinue the treatment because of the severity of the neuropathy and its major impact on their quality of life. Even though this study focused on initial discovery of association, we believe that this might be useful to suggest an alternative therapy for those patients before administration of vincristine. In this genome-wide association study, we identified several genetic loci associated with an incidence of VIPN ≥ grade 2. Based on the top-ranked SNPs in Table 6.1, 23 genes were identified in the association with either the primary or secondary neuropathy.

Among these 23 genes, one example is CSNK1G3. CSNK1G3 (Casein kinase 1, gamma 3) is an isoform of a monomeric serine-threonine protein kinase. Casein kinase has been implicated in a wide range of signaling activities such as cell differentiation, proliferation and apoptosis. Sakurai et al. previously found that the percentage of CK1ε-positive neurons and the expression level of CK1ε protein were increased in dorsal root ganglion and the spinal cord of the neuropathic mice (Sakurai et al., 2009). Another example is Nek6 (NIMA-related kinase 6). As RNA interference depletion studies reported by Yin et al., Nek6 and family isoforms are required for cell cycle progression through mitosis. Nek6 together with Nek7 and 9 form a mitotically activated module,
which plays a role in mitotic progression and more specifically spindle organization. Loss of Nek6 can lead to failure of centrosome separation in prophase and formation of weak mitotic spindles with reducing microtubule density (Yin et al., 2003). This modification of microtubule dynamic through the change of Nek6 expression can potentially lead to an association with neuropathy. In addition, MICAL3 is microtubule associated monooxygenase, calponin and LIM domain containing, which is an important regulator for microtubule polymerization (Terman et al., 2002; Fischer et al., 2005). Furthermore, ERCC8 is excision-repair-cross-complementing group 8. A study has shown that mutations in ERCC8 are associated with Cockayne syndrome, which is characterized by some neurological abnormalities including dys- or demyelinating neuropathy (Ohnishi et al., 1987; Smits et al., 1982).

Additional eight genes were discovered in an association with neuropathy from the gene enrichment analysis. In these eight genes, three (CACNAID, CACNA1C, CACNB1) are calcium channel, voltage-dependent subunits, and one (SCN8A) is a sodium channel, voltage gated subunit (Bock et al., 2011; Tan et al., 2011; Samak et al., 2011; Hori et al., 2012; Kloiber et al., 2012; Yang et al., 2012; Soeiro-de-Souza et al., 2012; Burgess et al., 1995). In particular, the phenotypes of mice with a mutation in SCN8A gene were found to be consistent with a defect in a neuronal sodium channel expressed in motor neuron (Burgess et al., 1995). Another two genes are in the transporter category. One (SLC29A4) is an equilibrative nucleoside transporter 4, which catalyzes the reuptake of monoamines into presynaptic neurons (Duan et al., 2013; Barnes et al., 2006). Another (SLC22A1) is a solute carrier family 22 member 1 which is a polyspecific organic cation transporter in many organs for elimination of endogenous small organic cations and toxins (Koehler et
al., 1997). In addition, two glutamate receptor, ionotropic subunits (GRIK1, GRIN3A) were also identified in the analysis. These two receptors have a critical role in excitatory synaptic transmission and plasticity in the CNS (Braga et al., 2009; Kaminski et al., 2004; Gryder et al., 2003; Rogawski et al., 2003). They govern a range of physiological conditions including neurological disorders caused by excitotoxic neuronal injury, psychiatric disorders and neuropathic pain syndromes (Petroff et al., 2002). These can be potential targets for further understanding of VIPN mechanism.

In the gene expression analyses, the expression levels were changed over 50% in eleven genes when treated with vincristine in the fibrosarcoma cell line. Interestingly, two (MICAL3 and ERCC8) and one (CANA1C) of these eleven genes were previously identified in the association and gene enrichment analyses, respectively. These findings confirmed that these genetic variants altered the gene expression, and the changes in expression may lead to toxicity. Different to the GEO analysis, two genes (COLEC11 and HSD18B4) found in the eQTL data were in a trans-association with the probe genes, and this indicated an indirect association with lipid biosynthesis. In fact, similar to what we observed in the pathway analysis, an association between VIPN and lipid biosynthesis mechanism was shown.

With the design of the analyses by using different association approaches, some direct or indirect associations between genomic variations and VIPN in children with ALL were identified. In the discovery phase of potential targets, we came across some similar correlations observed from different approaches which provided an additional confirmation of the findings. These findings could be clinical relevance given the widespread use of vincristine in treating childhood cancers. To validate these
observations, candidate genes will be sequenced, and the association between genomic variations and VIPN will be further examined in the sequencing data.
CHAPTERVII: Genetic signature to predict vincristine neuropathy and relapse in children with acute lymphoblastic leukemia

I. Introduction

A genome-wide association study (GWAS) is an examination of many common genetic variants in different individuals to determine if any variant is associated with a trait, and a way for scientists to identify genes involved in human disease. This method searches the genome for small variations that occur more frequently in people with a particular disease than in people without the disease. Each study can look at hundreds or thousands of single-nucleotide polymorphisms (SNPs) at the same time. Researchers use data from this type of study to pinpoint genes that may contribute to a person’s risk of developing a certain disease. Typically, GWAS focuses only on associations between SNPs and traits. There has been an increasing interest in relating genetic profiles to survival phenotypes such as time to event. Because of high dimensionality of SNP data, there is a limitation of collinearity in fitting a prediction model such as the Cox proportional hazards model (Sohn et al., 2009). To avoid the collinearity problem, several methods based on penalized Cox proportional hazards models have been proposed. One of the approaches that we used is a LASSO penalized Cox proportional hazard model (Tibshirani, 1997).

The benefit of using a LASSO approach, instead of other penalized regression models, is that this approach allows shrinking coefficients of insignificant predictors to exact zero which can simplify the model building in selecting a complex signature. In addition, this approach may potentially improve the predictability of genetic biomarkers to drug-induced toxicity as well as drug efficacy and provide a comprehensive and
unique signature. The objective of the study is to apply an alternative modeling approach to evaluate genetic biomarkers in predicting vincristine-induced neuropathy and relapse in childhood acute lymphoblastic leukemia patients. With this signature approach, a risk assessment of patients treated with vincristine can be further applied to distinguish the clinical outcome in a population based on the accuracy of the model performance and to provide clinically useful information to avoid unnecessary exposure to patients who are likely to experience severe toxicity but not benefit from the treatment.
Figure 7.1. **Model strategies using the LASSO penalized regression approach.**

Model 1: demographic and clinical variables were tested in the LASSO model; Model 2: demographic, clinical variables and SNPs with p-value < 1 \times 10^{-5} from the previous univariate analysis were tested in the LASSO model; Model 3: demographic, clinical variables and SNPs with p-value < 1 \times 10^{-4} from the previous univariate analysis were tested in the LASSO model.

Dom: dominant model; Rec: recessive model; Add: additive model.
2. Materials and Methods

2.1 Acute Lymphoblastic Leukemia (ALL) Patient Population and Study Design

Children with precursor B cell acute lymphoblastic leukemia from Pediatric Oncology Group (POG) 9904 and 9905 trials treated with 1.5mg/m²/dose vincristine doses and other chemotherapy treatments were included in the analysis as a training data set. The number of vincristine doses per patient over the course of treatment ranged from 18-23, depending on specific protocol and treatment arm. In this population, genotyping of germline samples from a total of 1068 subjects using an Affymatrix 6 assay remained in the analysis after a sequential data quality control including population stratification, sample and SNP missing call rate, Hardy-Weinberg disequilibrium and minor allele frequency. The details of data quality control were described in CHAPTER VI (Fareed et al., 2013). In addition, genotyping and clinical data of 122 ALL subjects from POG 9906 trial were used as a validation data set.

2.2 Phenotype Identification

As an extension to the association studies between genomic variations and vincristine-induced neurotoxicity in Chapter VI, this study continually evaluated and focused on both primary and secondary neuropathy as toxicity endpoints. The definition of the primary neuropathy based on National Cancer Institute Common Terminology Criteria for Adverse Events, version 3.0, is when patients experience sensor or motor neuropathy (grade ≥2). Time to event in this case means that the time to the first neuropathy event.

A secondary neuropathy which is considered as any sensory neuropathy, motor neuropathy or neuropathic pain ≥ grade 2 was also tested in the model. In addition to the
toxicity endpoints, an efficacy endpoint, the time to first relapse, was also taken into 
evaluation. By the definition of relapse of ALL patients, it included four types of relapse: 
an isolated bone marrow relapse: the presence of $\geq 25\%$ lymphoblasts in a bone marrow 
aspirate following the first complete remission (CR); CNS relapse: Positive 
cytomorphology and $> 5$ WBC/$\mu$L or positive cytomorphology with cerebrospinal 
fluid(CSF) WBC 0-4/$\mu$L on two successive occasions; testicular relapse: the histological 
evidence of lymphoblastic infiltration in one or both testes; and combined ALL relapse 
(protocol 9004; protocol 9905).

2.3 Genome-Wide Association Study

An association analysis of genotype frequencies with each phenotype was first 
carried out for the biomarker pre-selection process. Each phenotype was treated as a 
binary variable; ie, 1 for the event and 0 for no event. Clinical, demographic, and 
population stratification variables were also tested as covariates in the regression analyses. 
Genotype frequencies were evaluated in dominant, recessive and additive (gene-dose) 
models, and these analyses were performed in R coxph package (Therneau et al., 2014).

2.4 Genotype Imputation

To infer missing genotypes in the data, an imputation software called MACH (Li et 
al., 2009; Li et al., 2010) was applied to impute sporadic missingness of typed markers 
using a phasing approach with the following command: mach1 -d sample.dat -p 
sample.ped --states 200 -r 50 --dosage --prob --geno --phase After the imputation, the 
genotypes were coded as 0, 1 and 2 for minor allele copies.
2.5 LASSO Penalized Regression Model

After the association analysis in the GWAS study, each SNP was extracted with a statistical p-value and a hazard ratio (Therneau et al., 2014). Then, the SNPs were selected based on a pre-selection process which was defined in the modeling strategies (Figure 7.1). First, the smallest p-value across three genetic models was identified for each SNP. After ranking the p-value of each SNP, three following signature models were conducted. In model 1, only demographic and clinical variables were tested in the association. In model 2, in addition to variables tested in model 1, SNPs with a p-value < 1x10^{-5} were included in the association analysis. Furthermore, an extended SNP pool with p-value <1x10^{-4} were tested in model 3. The same modeling sequence was also examined with SNPs selected from only the dominant and additive models.

A particular feature of GWAS studies specifies that the dimension of the predictor space (number of genes) is typically larger than the number of samples (Sohn et al., 2009; Tibshirani et al., 1997; Benner et al., 2010; Fan et al., 2002). A penalized log partial likelihood in the LASSO method was utilized in conjunction with the proportional hazards model. The LASSO solution proposed by Tibshirani (Tibshirani et al., 1997) and Fan and Li (Fan et al., 2002) allows for parameter estimation by minimizing the log partial likelihood. The log partial likelihood was calculated using the following equation:

\[ l(\beta) = \sum_{i: c_i = 1} (\beta X_i - \log \sum_{j: y_j \geq y_i} \exp(\beta X_j)) \text{ subject to } \sum |\beta_j| \leq s \text{ where } s \text{ is a tuning parameter.} \]

If \( \sum |\beta_j| > s \), the regression coefficients are shrunken toward zero. In this context, imputed SNPs as well as clinical and demographic variables were analyzed in relation to the phenotype using the LASSO penalized Cox model in R package glmnet.
(Friedman et al., 2010). A cross-validation was implemented in the program to optimize lambda for model selection.

### 2.6 Model Evaluation

The predictive accuracy of a survival model was summarized using extensions of the proportion of variation explained by the model. The time-dependent sensitivity and specificity, and time-dependent receiver operating characteristic (ROC) curves were applied to evaluate the survival regression model by focusing on the correct classification rates. For survival data, there are several potential extensions of cross-sectional sensitivity and specificity. In this study, a cumulative/dynamic approach was performed to define the population who is classified as either a case or a control on the basis of vital status at time $t$ at any fixed time $t$. Each individual serves as a control for time $t < T_i$, but as a case for time $t \geq T_i$ (Heagerty et al., 2000; Heagerty et al., 2005). At any time point $t$, the sensitivity and specificity for a given threshold $c$ are defined as

**Sensitivity:** $P (M_i > c \mid T_i \leq t)$

**Specificity:** $P (M_i \leq c \mid T_i > t)$

Where $T_i$ is either time to event or censoring for the subject $i$, $M_i$ is a risk factor for the subject $i$. Furthermore, the nearest neighbor estimation (NNE) approach was applied to create a monotonic ROC curve. The sensitivity and specificity are based on joint and marginal distributions for time to event and risk factors, and the estimators of sensitivity and specificity were defined below

$$F_M(c) = \frac{\sum_{i=1}^{n} \mathbb{1}(M_i \leq c)}{n} \text{ where } \mathbb{1}(M_i \leq c) \text{ is the indicator function.}$$

$$S_{\lambda_n}(c, t) = \frac{1}{n} \sum_i^n S_{\lambda_n}(t \mid M_i = M_i) \mathbb{1}(M_i > c)$$
\[ \hat{S}_x(t|M = M_t) = \prod_{s \in r_n, s \leq t} \left\{ 1 - \frac{\sum_j 1(\hat{F}_M(M_j) - \hat{F}_M(M_{t})) \leq \lambda_n) 1(Z_j = s) \delta_j}{\sum_j 1(\hat{F}_M(M_j) - \hat{F}_M(M_{t})) \leq \lambda_n) 1(Z_j \geq s)} \right\} \]

with a chosen

\[ \lambda_n = n^{-1/3} \]

Where \( \hat{F}_M(c) \) is an estimator of the risk factor distribution; \( \hat{S}_x(c, t) \) is an estimator of the joint survival function; \( \hat{S}_x(t|M = M_t) \) is an estimator of the conditional survival function with a specific risk factor.

Then, the sensitivity and specificity were defined as

\[
\text{Sensitivity} = \frac{1 - F_M(c) - S_{\lambda_n}(c, t)}{1 - S_{\lambda_n}(-\infty, t)}
\]

\[
\text{Specificity} = 1 - \frac{S_{\lambda_n}(c, t)}{S_{\lambda_n}(-\infty, t)}
\]

### 2.7 Signature Models Validation

A signature for each phenotype constructed in the training data (POG 9904 and 9905 trials) was further validated in a completely independent validation dataset (POG 9906 trial) with a total of 122 ALL patients. To do so, SNPs which were identified in both training and validation data were first mapped. Among these selected SNPs and demographic variables, the estimated coefficient of each covariate from the univariate analysis in the training data were multiplied by the value of covariate in the validation data to generate a predictive risk score for each ALL patient in the validation data.

### 2.8 Clinical Application

To consider the clinical application, sensitivity and specificity calculated from ROC curves were evaluated to determine the cutoff for the risk score. Patients with a risk score less than the threshold were considered to be a low-risk group. In contrast, those who have a score above the threshold were assigned to be a high-risk group.
3. Results

To avoid the collinearity problem and high-dimensionality, a LASSO penalized Cox proportional hazard model was applied to the analysis. Different to a typical Cox proportional hazard model, the LASSO approach cannot handle missing genotypes which were observed commonly in the high-dimensional genome wide association study. In order to overcome this limitation, an imputation of typed SNPs was performed by MACH based on the haplotype of the dataset. At the initial stage of the model development, a signature model was introduced independently to test SNPs selected from each genetic model. As we moved forward to design a more comprehensive signature model, the strategy was modified and represented in Figure 7.1. For each phenotype, a minimum of p-value of each SNP was selected across either strategy 1, which included all three genetic models, or strategy 2, which only included the dominant and additive models. As Figures 7.2 A to L showed, the LASSO regression model in glmnet package allowed conducting a 10-fold cross-validation to optimize lambda which is a parameter to determine which coefficient was shrunken toward 0. The y-axis represented the partial likelihood error. The line on the right was drawn at the minimum error, and the other was drawn at the maximum value of lambda within 1 standard error of the minimum. At the maximum lambda, the LASSO coefficient paths were shown in Figures 7.2 M to X. The model was further evaluated by the ROC curves. This ROC analysis provided tools to select the optimal model, and the accuracy of the model was measured by the area under the ROC curve (AUC). The accuracy of the model represents how well the model separates the group being tested into those with and without the phenotype in question. For the primary neuropathy, no biomarker was selected in model 1 when only the
demographic and clinical factors were considered. The AUC is 0.7069419 in model 2, and the AUC increases to 0.9472682 in model 3. When the model was tested under the strategy of considering all three genetic models, the numbers of SNPs were increased in the test. As a result, the calculated AUCs improve to 0.908435 in model 2 and 0.9710504 in model 3, respectively. For the secondary neuropathy, the AUC is 0.5412067 without considering any genetic information. Furthermore, the AUCs increase to 0.7668192 and 0.9254674 in models 2 and 3, respectively. When the SNPs from the recessive model were also tested, the AUCs are 0.8876722 and 0.9691573 in models 2 and 3, respectively. The numbers of the selected predictors in each model were listed in Figures 7.3 E to F.

For relapse, the accuracy of the model was evaluated in a time dependent manner. As the duration of clinical follow-up lasted almost 10 years for some patients, the sensitivity and specificity of the model at four time points (1, 2, 4 and 8 years) were evaluated. In Figures 7.4 E to F, the AUCs are between 0.687354 and 0.712445 across four time points in model 1. Under the strategy 1, the AUCs are between 0.731131 and 0.756252 in model 2 and between 0.925097 and 0.933578 in the model 3 across time points. When additional SNPs selected from the recessive model were tested, the AUCs are between 0.837579 and 0.861887 in model 2 and between 0.926813 and 0.969303 in model 3. Overall, signature models with model 3 approach predicted both neuropathy and relapse more accurately than the models 1 and 2 based on the ROC curve evaluation.
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<th>Phenotype</th>
<th>Model Strategy and Model</th>
<th>Cross-validation</th>
<th>Lasso coefficient path</th>
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Figure 7.2. Cross-validation and LASSO coefficient path plots for primary neuropathy, secondary neuropathy and relapse. A-L represents the 10-fold cross-validation result with Model 2 and 3 as well as Dom/Rec/Add and Dom/Add genetic model strategies.
M-X represents the LASSO coefficient path result with Model 2 and 3 as well as Dom/Rec/Add and Dom/Add genetic model strategies. With the selected lambda, the coefficient of marker shrunk toward 0 on the right.
Dom: dominant model; Rec: recessive model; Add: additive model.
Furthermore, model 3 was used to identify patients who are at high risk of having an event based on selected sensitivity and specificity. A threshold of sensitivity at 0.90308144 and specificity at 0.901608868 were selected for primary neuropathy. The survival curves of high risk vs. low risk groups were plotted in Figures 7.5A and B. For the secondary neuropathy, sensitivity at 0.901325 and specificity at 0.900154 were selected to distinguish the survival of high risk vs. low risk groups in Figures 7.6A and B. Survival curves of high risk vs. low risk groups based on sensitivity at 0.900651992 and specificity at 0.9000682719 for relapse were shown in Figures 7.7A and B.

In order to evaluate the model predictability, the signature of each phenotype was further validated by predicting the risk score in a new population. There are 265,126 SNPs found in both training and validation sets. Among these SNPs, SNPs with p-value <1 x10^{-3} in the training set and p-value <1 x10^{-2} in the validation set were then evaluated. By taking the median risk score across the population, the time to primary neuropathy was compared between group 1 (>median risk score) and group 2 (<median risk score). A p-value of 0.00066 was given based on $\chi^2$ distribution in the log rank test. Similarly, a significant p-value of 0.00791 in $\chi^2$ distribution was observed when comparing the time to secondary neuropathy between group 1 (>median risk score) and group 2 (<median risk score). For the efficacy endpoint, the difference of time to relapse between the two groups, based on a cutoff of the median predictive risk score from the validation data, was with a p-value of 0.000656. All these findings indicated that the signature of each phenotype was well validated in an independent population based on the proposed validation scenario.
In addition, to apply the clinical utility of the signature model by identifying patients who are less likely to be benefited from the treatment, patients at high risk of relapse were selected by a risk score greater than 222.009734 based on specificity at 0.9000682719. Furthermore, a subgroup of those patients who are likely to experience primary neuropathy was identified based on a risk score greater than 1 at sensitivity of 0.820777. A total of 48 individuals were identified in this category (Figure 7.8). A sub-population who is most likely to receive the benefit of the treatment was also classified in Figure 7.8. Similar to the primary neuropathy, patients were divided into four sub-groups based on the risk assessment of both relapse and secondary neuropathy in Figure 7.9.
### Phenotype | ROC curve | Model strategy | Biomarkers
---|---|---|---
Primary neuropathy | ![A](image1.png) | Dom/Rec/Add | E
  - 1: 6 | 0 | Dem and SNPs < 10E-5
  - 2: 70 | 35 | Dem and SNPs < 10E-5
  - 3: 289 | 132 | Dem and SNPs < 10E-4
Primary neuropathy | ![B](image2.png) | Dom/Add | F
  - 1: 4 | 0 | Dem
  - 2: 20 | 8 | Dem and SNPs < 10E-5
  - 3: 107 | 51 | Dem and SNPs < 10E-4
Secondary Neuropathy | ![C](image3.png) | Dom/Rec/Add | G
  - 1: 4 | 1 | Dem
  - 2: 74 | 48 | Dem and SNPs < 10E-5
  - 3: 291 | 140 | Dem and SNPs < 10E-4
Secondary Neuropathy | ![D](image4.png) | Dom/Add | H
  - 1: 4 | 3 | Dem
  - 2: 30 | 12 | Dem and SNPs < 10E-5
  - 3: 126 | 50 | Dem and SNPs < 10E-4
Figure 7.3. **ROC curves of primary and secondary neuropathy in two genetic model strategies and the numbers of tested and selected biomarkers in the models.**

Dom: dominant model; Rec: recessive model; Add: additive model; Dem: demographic and clinical factors.
**Figure 7.4. Time dependent ROC of relapse in two genetic models.**

A-B: ROC curves from 1 to 8 years

C-D: The number of tested and selected biomarkers in model 1 to 3.

E-F: Area under the ROC curve (AUC) of each model at time 1, 2, 4 and 8 years.

Dom: dominant model; Rec: recessive model; Add: additive model.
Figure 7.5. Survival plot of high risk vs. low risk group of primary neuropathy with (A) sensitivity=0.903081 and (B) specificity=0.901608. (A) Patients have risk score greater than 9.15 considered as high risk group; (B) Patients have risk score greater than 7.26 considered as high risk group.
Figure 7.6. **Survival plot of high risk vs. low risk group of secondary neuropathy** with (A) sensitivity=0.901325 and (B) specificity=0.900154.

(A) Patients have risk score greater than 8.01 considered as high risk group; (B) Patients have risk score greater than 6.34 considered as high risk group.
Figure 7.7. Survival plot of high risk vs. low risk group of relapse with (A) sensitivity=0.900651 and (B) specificity=0.900068.
(A) Patients have risk score greater than 204.74 considered as high risk group; (B) Patients have risk score greater than 222.01 considered as high risk group.
Figure 7.8. A graphical representation of the clinical application using the signature model to identify ALL patient in four subgroups (HR RE/HR NE; HR RE/ LR NE; LR RE/HR NE; LR RE/ LR NE) based on relapse and primary neuropathy endpoints and their clinical recommendation.

NE: primary neuropathy; RE: relapse; HR: high risk; LR: low risk.
Figure 7.9. A graphical representation of the clinical application using the signature model to identify ALL patient in four subgroups (HR RE/HR NEP; HR RE/LR NEP; LR RE/HR NEP; LR RE/LR NEP) based on relapse and secondary neuropathy endpoints and clinical recommendation.

NEP: secondary neuropathy; RE: relapse; HR: high risk; LR: low risk
4. Discussion

Our analysis of single nucleotide polymorphism data demonstrated that the LASSO penalized regression is capable of identifying pertinent predictors. The LASSO technique for variable selection in the Cox model seemed to be a worthy competitor to stepwise selection. It is less variable than the stepwise approach and still yields interpretable models. The computational time is relatively quick compared to the traditional GWAS study. Initially, we expected that the model would not only be capable of predicting time to event by a genetic pattern, but also would provide a reasonable prediction in a relative small association data set with no statistical significant association identified in the GWAS study. Based on our study design, we constructed three models to compare how much information would be enough in the test to generate a reasonable prediction to the same phenotype. As the model performed 10-fold cross-validation, it allowed the function to run 10fold+1 times to get the lambda sequence and compute the fit with each of the folds omitted. The optimal lambda was selected based on the mean of accumulated cross-validated error. This process was done repeatedly in each model and generated a unique signature for each phenotype.

Based on ROC curve evaluation, the area under the curve of ROCs in model 3, which contained demographic and SNPs with p-value <1x10^{-4} across both neuropathy and relapse, generally exceeded 0.9. This indicated that the model well predicted patients with and without event. All important findings are subject to replication. This attitude allows us to define the most informative SNPs rather than only declaring their global significance. Our approach to data analysis is motivated by this consideration. In addition, the signature of each phenotype was further validated in the validation data set. In this
backward validation, the overlapped SNPs with a predefined p-value threshold were first identified; then, the predicted risk scores were calculated for each individual in the validation data using estimated coefficients from univariate analysis in the training data. As the result indicated a statistically significant separation of patients’ time to neuropathy or relapse based on the median of predicted risk score using the log-rank test, this validation gave a preliminary overlook of how well the model predicts based on the best scenario. However, the current validation data set, which mainly has a population at high risk for treatment failure, may not be the best candidate to validate the signature model. Therefore, further analysis needs to be done, not only to validate the model by taking all the tested SNPs into consideration which may be more precise in evaluating the signature applicability, but also to evaluate the signature in a standard risk disease population similar to POG 9904 and 9905 trials.

In addition to validating the generalizability of the model, the clinical application of the signature model is also a major focus in this study. Clinically, as ALL relapse still carries a very poor prognosis and the majority of relapse continues to occur in patients without apparent “high-risk” features (Stanulla et al., 2009; Schrappe et al., 2000a; Schrappe et al, 2000b), the utility of the signature model becomes meaningful when a subgroup of patients who have no benefit from the treatment because of the high risk of relapse and neurotoxicity can be predicted and alternative treatments can be recommended. Similarly, for those who are either at high risk of relapse or high risk of neuropathy, the clinicians can evaluate the cost and effectiveness to recommend dose modification. Those who are less likely to experience toxicity or relapse can be the best targets for the treatment and for understanding drug exposure variability. With all these
factors taken into account, our signature modeling approach not only provides an innovated assessment to high dimensional genomic data for identifying potential associations with toxicity and efficacy, but also integrates a statistical model with clinical useful application and interpretation.
CHAPTER VIII: Conclusions and future directions

1. Conclusions

The major foci of this dissertation were to explore the impact of modeling and simulation in drug development and patient care: 1) where a population approach in pharmacometrics can be used to understand pharmacokinetics of drugs in humans from a translational in vivo model, the covariate effect on exposure variability in a special population, the pharmacodynamics of biomarkers for predicting the clinical outcome of a treatment and the evaluation of a trial design based on a simulated virtual population; and 2) how a statistical modeling approach can be utilized to evaluate the association between pharmacogenomic variations, drug-induced toxicity and efficacy to establish a signature pattern that determines patients’ risk of experiencing these effects. The analyses from this dissertation address limitations of each modeling approach presented. These limitations are tied back to the underlying assumptions in the modeling approach as well as the potential sources of variability that can contribute to trial failure and should be accurately measured. This dissertation also points out an integration between different modeling approaches which potentially allows more sophisticated understanding of the biomarker application and helps to capture disease status and treatment effect across a spectrum of modeling approaches and data types.

Based on the major findings outlined above, the overarching conclusion of each modeling and simulation application can be drawn concerning the body of work presented in this dissertation. These are: 1) a translational PK modeling approach that was able to predict clozapine and norclozapine CNS exposures in humans relating these exposures to receptor binding kinetics at multiple receptors; 2) a population
pharmacokinetic analysis of a study of sertraline in depressed elderly patients with Alzheimer’s disease (DIADS-2) identified site specific differences in drug exposure contributing to the overall variability in sertraline exposure; 3) the utility of a longitudinal tumor dynamic model developed by the Food and Drug Administration for predicting survival in NSCLC patients drawn from NCI funded studies, including an exploration of the limitations of this approach; 4) a Monte Carlo clinical trial simulation approach that was used to evaluate a pre-defined oncology trial with a sparse drug concentration sampling schedule with the aim to quantify how well individual drug exposures, random variability, and the food effects of abiraterone and nilotinib were determined under these conditions; 5) a time to event analysis that facilitated the identification of candidate genes including polymorphisms associated with vincristine-induced neuropathy from several association analyses in childhood ALL patients; and 6) a LASSO penalized regression model that predicted vincristine-induced neuropathy and relapse in ALL patients and provided the basis for a risk assessment of the population. Each of these conclusions is summarized followed by further directions in the section which follows.

A. A translational PK modeling approach that predicted clozapine and norclozapine CNS exposures in humans relating these exposures to receptor binding kinetics at multiple receptors

From a clinical perspective, it is important to understand the broad receptor coverage of clozapine that is thought to account for clozapine’s unique superiority in treatment resistant schizophrenia. With the innovation and improvement of measurement techniques, clozapine concentrations now can be measured at the target site using microdialysis so that the concentration can be more precisely related to response at the
effect site (Derendorf et al., 2000; Chefer et al., 2009; Jacobson et al., 2013). As part of this dissertation (CHAPTER II), we demonstrated that the proposed PK model can be extrapolated to predict human systemic exposure including clozapine brain concentrations and associated receptor occupancy profiles in humans at clinically relevant doses. In addition, the model simultaneously captured the parent-drug and metabolite in the system. This is relevant as norclozapine also has activity at multiple receptors in the CNS that could contribute to response (Bishara et al., 2008). This modeling approach could form the foundation for the design of the future comprehensive PK-PD models as well as extend our understanding of clozapine’s complex behavioral effects in humans. Furthermore, this model framework can serve to support the discovery of a new drug entity that may share the same mechanism of action as clozapine.

B. A population pharmacokinetic analysis of a study of sertraline in depressed elderly patients with Alzheimer’s disease (DIADS-2) identified site specific differences in drug exposure contributing to the overall variability in sertraline exposure

This is the first population pharmacokinetic (PPK) study of sertraline focusing on AD patients with depression. Compared to younger patients, the elderly have a longer half-life, but there was no difference between elderly healthy volunteers in the literature and these AD patients in PK characteristics. In addition, a covariate analysis was conducted to explore whether the variability of the PK parameter estimates could be explained by patient specific characteristics. Based on the analysis, the clinical site was the only significant covariate explaining variability in the CL/F change. The clinical implications for subjects at the outlying site would be important if this represented a true bias in subject selection. As we discussed in CHAPTER III, no significant difference in
demographic or clinical attributes between this study site and the others were found; therefore, a potential source of variation is the implementation of study procedures at this site. In conclusion, site variability in trial procedures is critical to the validity of multi-site trials, and population pharmacokinetics may prove helpful in this assessment. With the application of the population PK model approach, a close monitoring of dosage regimen and administration is recommended in the future multicenter study especially with sparse PK sample collections so that patients would not exposed to unnecessary harm. This can also minimize what may be artificial systematic differences in exposure, and therefore potentially confound on the interpretation of response and/or toxicity.

C. The utility of a longitudinal tumor dynamic model developed by the Food and Drug Administration for predicting survival in NSCLC patients drawn from NCI funded studies, including an exploration of the limitations of this approach

Previous studies have shown that modeling and simulation is an efficient means to support decision making at the Phase 2 to Phase 3 transition in drug development (Claret et al., 2012), therefore, it was thought that this longitudinal tumor model could serve as a framework to predict clinical outcomes of cancer patients ideally resulting in smaller trials with shorter observation periods (Adjei et al., 2009; Dhani et al., 2009; Maitland et al., 2011; Maitland et al., 2011; Mandrekar et al., 2010; Stein et al., 2011; Yap et al., 2010). Based on this initial idea, we demonstrated in CHAPTER IV that a modeling strategy to assess the tumor load dynamics found that the tumor model parameter estimates of tumor shrinkage and progression rates were both lower than the findings from the previous FDA large registration trial data. However, when the FDA trial was reduced to a similar size as our CALGB30203 study and patient characteristics matched,
the parameter estimates became comparable. The lower parameter values for tumor dynamics from the model using the data from the CALGB trials suggested that the prediction of overall survival using this metric was likely impacted by the sample size and heterogeneity in the tumor presentation. In addition, a simplified time to tumor growth (TTG) parameter was observed to be more robust in predicting overall survival in our small trial as a single factor, but the clinical interpretation and utility need to be further explored. In conclusion, the data quality is critical for a model to give a meaningful contribution to predicting patient outcome. The limitations inherent in tumor measurement can lead to the misrepresentation of the true tumor growth dynamic. This variation can be more problematic when the sample size is small. Further analysis is required to explore the utility of the tumor dynamic model in predicting patients’ survival outcomes using alternative measurements such as automated capture of the three-dimensional tumor volumes from raw CT scan data (Schwartz et al., 2000). It appears that this approach may be more accurate in describing the tumor change over time, and potentially eliminates the inter-rater variability from the traditional sum of the longest diameter measurement carried out by radiologists. In addition, a simulation approach could be applicable to evaluate how different level of variance of the tumor measurements can affect the overall tumor model parameter estimates. Until more precise and/or accurate measures are available, it will also be difficult to ascertain the true relationship between tumor loads (especially across tumor types that may be treated with agents that are –static instead of –cidal in their effects) and patient specific outcomes such as progression free survival and overall survival.
D. A Monte Carlo clinical trial simulation approach was used to evaluate a pre-defined oncology trial with a sparse drug concentration sampling schedule with the aim to quantify how well individual drug exposures, random variability, and the food effects of abiraterone and nilotinib were determined under these conditions.

Food intake is one important contributor to variability in oral bioavailability (Singh et al., 2004; Winstanley et al., 1989; Gu et al., 2007) in particular for some of the newer oral anticancer agents such as abiraterone and nilotinib (Ryan et al., 2010; Yin et al., 2010). This can cause a significant impact on systemic drug exposures that can lead to drug toxicity and/or therapeutic failure (Koch et al., 2009; Kang et al., 2010). We demonstrated in CHAPTER V how a clinical trial simulation using a Monte Carlo method under the framework of a Pop PK approach provided understanding of a clinical oncology population pharmacokinetic trial design with a random sampling schema. More specifically, we were able to quantify how well this design and the data collection scheme associated with it and were able to capture the variability and the magnitude of individual drug exposure in the presence of a large food effect in nilotinib and abiraterone trials. This simulated approach facilitated an early evaluation of the proposed trial design. However, clinical trial simulations are generated based on many trial assumptions, and these assumptions may include uncertainty. For future studies, different scenarios with underlying assumptions can be tested to optimize the design. With the limitations of current analysis discussed in CHAPTER V, additional components such as a drop-out pattern of the trial and patient compliance, which may vary in the clinical practice setting, should be explored in future analyses.
This study emphasizes the importance of addressing trial designs where intensive sampling cannot be obtained and yet there is a need to understand the drug exposure characteristics to what are otherwise medications with highly variable pharmacokinetic dispositions.

E. A **time to event analysis facilitated the identification of candidate genes including polymorphisms associated with vincristine-induced neuropathy from several association analyses in childhood ALL patients**

Vincristine is among the most commonly used anticancer agents, however, little is known regarding its disposition and optimal dosing. This gap in knowledge can lead to negative clinical outcomes such as toxicity due to drug overdosing or lack of efficacy due to sub-therapeutic dosing. Vincristine is associated with highly variable cumulative dose-dependent peripheral neuropathy. To explore the association between germline variants and VIPN in pediatric ALL patients, a genome-wide association analysis was carried out initially. The analysis was expanded to examine the association using gene enrichment analysis. In both analyses, we identified several genetic loci associated with incidence of \( \geq \) grade 2 VIPN. Based on the top-ranked SNPs, 31 genes were found in an association with either primary or secondary neuropathy. In **CHAPTER VI**, a discussion was carried out to identify those candidate genes and their corresponding pathways. In this stage of discovery, we found that there are many genes which are indirectly associated with the mechanisms of leading vincristine-induced neuropathy, but we believe these findings can be potential focuses for the future *in vitro* studies after association is confirmed by sequencing an independent data set from a multi-center trial. Furthermore, the candidate genes selected from the association analysis were evaluated in the
interactive pathway analysis and gene expression analysis. Based on both the GEO and eQTL databases, few additional gene candidates appeared in association with neuropathy. In conclusion, we came across some similar correlations observed from different associations and data approaches that provided additional confirmation of the findings. These findings have significant potential clinical relevance given the widespread use of vincristine in treating childhood cancers.

F. A LASSO penalized regression model predicted vincristine-induced neuropathy and relapse in ALL patients and provided the basis for a risk assessment of the population

As the previous section (CHAPTER VI) discovered the association between germline variants and VIPN using multiple association analyses, this chapter focused on an alternative modeling approach to identify a genetic signature pattern which allows predicting the traits and avoid the collinearity in fitting which is a common limitation found in typical Cox proportional hazards model (Sohn et al., 2009). By performing a LASSO penalized regression model, a signature was developed not only for neuropathy but also for relapse independently as previous study showed no correlation between two phenotypes. Under different model strategies, we demonstrated that a unique genetic signature was capable to predict neuropathy or relapse with reasonable precision based on ROC evaluations. In order to indicate the clinical utility of the signature model, a risk assessment was carried out under the structure of genetic signature. In sum, the whole population was divided into four major groups based on the statistical threshold for both relapse and neuropathy. This identification would provide clinicians an initial guidance for dose recommendations which might result in a more efficient treatment plan and avoid unnecessary drug exposure to patients. The utility of the signature model seems
promising in predicting time to event, but a more comprehensive validation plan will need to further explore and test the model predictability. In fact, instead of building a signature only based on typed SNPs from the genotyping data, an additional approach was suggested to expand the signature by examining the untyped SNPs from the imputation. This approach might provide a more representative pattern of predictive signature across different populations and data, but the quality of imputation data needs to be carefully evaluated as we found to be a challenge initially when we imputed our training data. In terms of clinical utility of the signature, a challenge we are facing right now is if there is a surrogate endpoint which can help us to understand the disease and treatment effect as early as possible and enhance the application of the modeling approach. To consider this, minimal residual disease which was suggested by some studies (Campana 2010; Borowitz MJ, et al., 2008) as an important predictor of ALL relapse may be valuable to explore in this case.

Overall, this dissertation demonstrated the utility of modeling and simulation in various study design scenarios and its application and contribution for the future studies in drug development and patient care. Although different population and disease were focused in each chapter to explore a specific underlying modeling approach, the whole dissertation provided a projection of how each approach can be integrated in drug development process. An integration of multiple modeling approaches is also suggested by this dissertation to overcome the challenges and to examine the proposed solutions for the limitations of the current modeling approach. As recalled the drug development process in Figure 1.1, if there is a new compound targeted at early drug discovery phase, a combination of preclinical and translational modeling approach (CHAPTER II) can
capture its exposure-response relationship *in vivo* system, extrapolate a first to human dose and further predict some pharmacological relationships which have not been assessable in the human system. As this new compound is moved forward to the clinical phase, PPK model (CHAPTER III) can serve as a tool to continuously evaluate its exposure-response relationship, quantify the level of variability and identify covariate effect. This approach can be further expanded to predict PK in a special population such as elderly or another disease population. In addition to understand the PK profile of this new compound, assessment of the dynamics of a biomarker (CHAPTER IV, VI and VII) can provide insight into the disease progression and capture a therapeutic drug effect of this compound which might be useful for early go/no go decision or personalized medicine later on before the clinical endpoints are available. Meanwhile, a clinical trial simulation (CHAPTER V) can explore various dosage regimens of the compound and optimize the trial design which might allow reducing the numbers of the studies and be more efficient and informative in bridging studies from one phase to another. As a demonstration of how these approaches emphasized in the dissertation can be integrated in the drug development process, we also need to be aware that there are many more assessments could be done by modeling and simulation in the process.

In summary, modeling and simulation is a tool that enhances our understanding of pharmacokinetics and pharmacodynamics for specific drug entities and provides critical recommendations for dose selection and treatment plans. Leveraging the findings from biological studies, it will continue to be an essential element in drug development and patient care.
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EDUCATION TRAINING

Doctor of Philosophy (Medical and Molecular Genetics) 2009-08/2014
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Disease and Therapeutic modeling program Training 2010-2013
2010: R Programming for Pharmacometrics (Metrum Institute)
2010: PHAR-F 813 Clinical Pharmacokinetics (Indiana University)
2011: Monolix 3.2 workshop (INRIA, ACOP)
2011: New and Advanced Features of NONMEM7&PDx-Pop4
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2011: Uppsala Pharmacometric Summer School
2011: PHAR-F 836 Pharmacogenetics and Physiologic Disposition of Drugs (Indiana University)
2011: Introduction to Bayesian PK-PD modeling & simulation (Metrum Institute)
2012: Pharmacokinetic-pharmacodynamic modeling (State University of New York at Buffalo)
2012: 2-Day workshop on monoclonal antibody pharmacokinetics & Pharmacodynamics (State University of New York at Buffalo)
2012: PKPD-modeling of Continuous and Categorical data in NONMEM (Uppsala University, PAGE)
2013: Quantitative System Pharmacology (ASCPT)
2013: New & Advanced Methods of NONMEM 7 & PDx-Pop 5.0 including New Features in the Upcoming Release of NONMEM 7.3 (ICON Development Solutions, ACOP)
2013: Using MATLAB® and SimBiology® for Mechanistic Systems Pharmacology Workflows (Mathworks, ACOP)
2013: Pharmacokinetic-pharmacodynamic modeling from Sunny Buffalo (Genentech)
PROFESSIONAL EXPERIENCES

June 2013-Sep 2013  **Genentech**  [Internship]
Clinical Pharmacology, Genentech, South San Francisco, CA

May 2008-Jul 2009  **Volt Temp Services at Eli Lilly and Company**  [Biochemist-Contractor]
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RESEARCH EXPERIENCES

2013-2014  **Population pharmacokinetic modeling of vincristine and metabolite in children with acute lymphoblastic leukemia from Children’s Oncology Group**
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Mentor: Robert R. Bies, PharmD, Ph.D.

2013  **Pharmacokinetic and pharmcodynamic modeling of antibody-drug conjugate target glycosylated antigen in patients with ovarian cancer**
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2012-2014  **Develop a genetic signature model to predict efficacy and toxicity using GWAS analysis by Lasso penalized logistic regression in Childhood Acute Lymphoblastic Leukemia patients**
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2012-2013  **Prediction of brain clozapine and norclozapine concentrations in human using a scaled PK model for rat brain and plasma PK**
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2012-2013 Limited on longitudinal tumor size modeling of non-small cell lung cancer with data collected from completed cooperative group trials
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2009-2010 Population pharmacokinetic modeling of ziprasidone in patients with schizophrenia from the CATIE study
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Mentor: Yingying Guo, Ph.D.

2008 Neo gene construction into an Escherichia coli cell.
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TEACHING EXPERIENCES (LECTURE)

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TEACHING ASSISTANT EXPERIENCES

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SYMPOSIUM HOSTING EXPERIENCES

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2012  The Second Indiana Clinical and Translational Sciences Institute (CTSI)
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2011  The First Indiana Clinical and Translational Sciences Institute (CTSI)
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Awards/Honors/Citations
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PUBLICATIONS
1. Li CH, Li L, Zhang P, Renbarger JL. Genetic signature to predict vincristine neuropathy and relapse in children with acute lymphoblastic leukemia. [In preparation]

2. Li CH, Li L, Skiles JL, Then, JE, Renbarger JL. Genetic determinants of vincristine neuropathy in pediatric acute lymphoblastic leukemia patients. [In preparation]


6. Li CH, Sherer EA, Lewis LD, Bies RR. Clinical trial simulation to evaluate study design: capturing abiraterone and nilotinib exposure. [In preparation for JCP]


ABSTRACTS AND SCIENTIFIC PRESENTATIONS


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**PROFESSIONAL ASSOCIATION MEMBERSHIPS**

2012-Present  **American Association of Pharmaceutical Scientist (AAPS)**

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**SCIENTIFIC MEETINGS**

2011  **American Conference on Pharmacometrics (ACOP)** San Diego, CA

2011  **Alliance Annual meeting (Cancer Leukemia Group B)** Chicago, IL

2011  **The First Indiana Clinical and Translational Sciences Institute (CTSI) Symposium on Disease and Therapeutic Response Modeling** Indianapolis, IN

2012  **Population Approach Group in Europe (PAGE)** Venice, Italy

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