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Preclinical Pharmacology of the MDM2 Antagonist Nutlin-3a

Fan Zhang
University of Tennessee Health Science Center

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PRECLINICAL PHARMACOLOGY OF THE MDM2 ANTAGONIST
NUTLIN-3A

A Dissertation
Presented for
The Graduate Studies Council
The University of Tennessee
Health Science Center

In Partial Fulfillment
Of the Requirements for the Degree
Doctor of Philosophy
From The University of Tennessee

By
Fan Zhang
December 2011
DEDICATION

I dedicate this dissertation to my parents,
    Hua Zhang and Yufang Zan,
    and to my husband and son,
    Fei Ma and Alexander Zhang Ma.
ACKNOWLEDGEMENTS

I would like to take this opportunity to express my deepest gratitude to all those who supported and helped me during my research. First and foremost, I would like to thank my mentor, Dr. Clinton F. Stewart, for his guidance, support, encouragement, and inspiration. I am incredibly lucky to be able to be a student of Dr. Stewart and to learn from his wisdom. I sincerely appreciate Dr. Stewart for everything that he has taught me.

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In addition, I want to thank every member and friend in the Stewart laboratory. Thanks very much to our lab supervisor Dr. Stacy Throm, former supervisor Laura Miller, and co-workers Dr. Laura Lea Murley, Dr. Michael Tagen, Dr. John Panetta, Dr. Fan Wang, Zaifang Huang, Dr. Feng Bai, Mohamed Elmegiey, Dr. Steven Zatechka, Thandranese Owens, Jenkin Chen, Daniel Groepper, and Rachel Kennedy for their help throughout the project. In addition, I would also like to thank Jeremy Mallari, Katie Nemeth, Fangyi Zhu, Jiakun Zhang, Min Lu, Nidal Boulos, Lei Yang, Shelly Wilkerson, and Frederique Zindy for their help in the nutlin-3a pharmacokinetic study.

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Finally, I extend deep thanks to my parents, my husband, and my parents -in-law for their love and support.
Nutlin-3a is an MDM2-p53 interaction antagonist that is under investigation in preclinical models for a variety of pediatric malignancies, including neuroblastoma, retinoblastoma, leukemia, and rhabdomyosarcoma. In the current research, we conducted preclinical pharmacology studies of nutlin-3a to evaluate the synergistic effect of the nutlin-3a and topotecan combination on neuroblastoma cell growth, to assess the effect of nutlin-3a on breast cancer resistance protein (BCRP), and to characterize the disposition of nutlin-3a in the mouse plasma and multiple tissues.

Activating the p53 pathway might offer a new therapy for neuroblastoma. In the first part of the study, we assessed the effect of nutlin-3a on the cell viability of neuroblastoma both as a single agent and in combination with topotecan. We showed that targeting MDM2-p53 interaction using nutlin-3a reduced cell growth in neuroblastoma cells. p53 wild-type cells were much more sensitive to nutlin-3a treatment compared to p53 mutant cells. When nutlin-3a was combined with topotecan, a synergistic effect on neuroblastoma cell growth was observed. To explore the mechanism of synergy, we performed quantitative real-time polymerase chain reaction (qRT-PCR) and western blot analysis and found reduction of P-gp expression at both the message level and protein level in p53 wild-type neuroblastoma cells. This is the first study showing the synergistic effect of nutlin-3a in combination with topotecan in neuroblastoma cells and the reduction of P-gp expression by nutlin-3a in p53 wild-type cells.

Although nutlin-3a is currently under pre-clinical investigation as a p53 reactivation agent, it has been recently demonstrated also to have p53 independent actions in cancer cells. In the second part of the study, we first reported that nutlin-3a can inhibit the efflux function of BCRP. We observed that although the nutlin-3a IC₅₀ did not differ between BCRP over-expressing and vector control cells, nutlin-3a treatment significantly potentiated the cells to treatment with the BCRP substrate mitoxantrone. Combination index calculations suggested synergism between nutlin-3a and mitoxantrone in cell lines over-expressing BCRP. Upon further investigation, it was confirmed that nutlin-3a increased the intracellular accumulation of BCRP substrates such as mitoxantrone and Hoechst 33342 in cells expressing functional BCRP without altering the expression level or localization of BCRP. Interestingly, nutlin-3b, considered virtually "inactive" in disrupting the MDM2/p53 interaction, reversed Hoechst 33342 efflux with the same potency as nutlin-3a. Intracellular accumulation and bi-directional transport studies using MDCKII cells suggested that nutlin-3a is not a substrate of BCRP. Additionally, an ATPase assay using Sf9 insect cell membranes over-expressing wild-type BCRP indicated that nutlin-3a inhibits BCRP ATPase activity in a dose-dependent fashion. In conclusion, our studies demonstrate that nutlin-3a inhibits BCRP efflux function, which consequently reverses BCRP-related drug resistance.

Understanding drug disposition is critical in preclinical drug development. In the third part of the study, we used physiologically-based pharmacokinetic (PBPK) modeling to characterize the disposition of nutlin-3a in mice. Plasma protein binding and blood
partitioning were assessed by \textit{in vitro} studies. After intravenous (10 and 20 mg/kg) and oral (50, 100, and 200 mg/kg) dosing, tissue concentrations of nutlin-3a were determined in plasma, liver, spleen, intestine, muscle, lung, adipose, bone marrow, adrenal gland, brain, retina, and vitreous fluid. The PBPK model was simultaneously fit to all pharmacokinetic data using NONMEM. Nutlin-3a exhibited nonlinear binding to murine plasma proteins, with the unbound fraction ranging from 0.7 to 11.8%. Nutlin-3a disposition was characterized by rapid absorption with peak plasma concentrations at approximately 2 h and biphasic elimination consistent with a saturable clearance process. The final PBPK model successfully described the plasma and tissue disposition of nutlin-3a. Simulations suggested high bioavailability, rapid attainment of steady state, and little accumulation when administered once or twice daily at dosages up to 400 mg/kg. The final model was used to perform simulations of unbound tissue concentrations to determine which dosing regimens are appropriate for preclinical models of several pediatric malignancies.

In conclusion, our results showed that nutlin-3a synergistically inhibited the growth of neuroblastoma cells when combined with topotecan. Nutlin-3a reversed BCRP-mediated drug resistance by inhibiting the function of BCRP. A PBPK model was successfully established to describe the disposition of nutlin-3a in plasma and tissues of interest for pediatric malignancies.
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<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>ABC</td>
<td>ATP-binding cassette</td>
</tr>
<tr>
<td>ADME</td>
<td>Adsorption, distribution, metabolism, and excretion</td>
</tr>
<tr>
<td>ADP</td>
<td>Adenosine diphosphate</td>
</tr>
<tr>
<td>AIC</td>
<td>Akaike's information criterion</td>
</tr>
<tr>
<td>ALL</td>
<td>Acute lymphoblastic leukemia</td>
</tr>
<tr>
<td>AML</td>
<td>Acute myelogenous leukemia</td>
</tr>
<tr>
<td>ATP</td>
<td>Triphosphate</td>
</tr>
<tr>
<td>AUC</td>
<td>Area under the curve</td>
</tr>
<tr>
<td>BBB</td>
<td>Blood-brain barrier</td>
</tr>
<tr>
<td>BCEC</td>
<td>Brain capillary endothelial cells</td>
</tr>
<tr>
<td>BCRP</td>
<td>Breast cancer resistance protein</td>
</tr>
<tr>
<td>BCS</td>
<td>Biopharmaceutics classification system</td>
</tr>
<tr>
<td>BDDCS</td>
<td>Biopharmaceutics drug disposition classification system</td>
</tr>
<tr>
<td>CI</td>
<td>Combination index</td>
</tr>
<tr>
<td>CL</td>
<td>Clearance</td>
</tr>
<tr>
<td>CNS</td>
<td>Central nervous system</td>
</tr>
<tr>
<td>CSF</td>
<td>Cerebrospinal fluid</td>
</tr>
<tr>
<td>DDI</td>
<td>Drug–drug interaction</td>
</tr>
<tr>
<td>DMEM</td>
<td>Dulbecco’s modified eagle’s medium</td>
</tr>
<tr>
<td>DMSO</td>
<td>Dimethyl sulfoxide</td>
</tr>
<tr>
<td>Fa</td>
<td>Fraction of absorption</td>
</tr>
<tr>
<td>FBS</td>
<td>Fetal bovine serum</td>
</tr>
<tr>
<td>FDA</td>
<td>Food and drug administration</td>
</tr>
<tr>
<td>Fg</td>
<td>Fraction of intact drug escaping gut metabolism</td>
</tr>
<tr>
<td>GST</td>
<td>Glutathione S-transferase</td>
</tr>
<tr>
<td>HBSS</td>
<td>Hank’s balance salt solution</td>
</tr>
<tr>
<td>IND</td>
<td>Investigational new drugs</td>
</tr>
<tr>
<td>LLOQ</td>
<td>Lower limit of quantification</td>
</tr>
<tr>
<td>MDCK</td>
<td>Madin-Darby canine kidney epithelial</td>
</tr>
<tr>
<td>MDM2</td>
<td>Murine double minute-2</td>
</tr>
<tr>
<td>MDR</td>
<td>Multiple drug resistance</td>
</tr>
<tr>
<td>MRP1</td>
<td>Multidrug resistance protein 1</td>
</tr>
<tr>
<td>MXR</td>
<td>Mitoxantrone resistance protein</td>
</tr>
<tr>
<td>NB</td>
<td>Neuroblastoma</td>
</tr>
<tr>
<td>NBD</td>
<td>Nucleotide binding domain</td>
</tr>
<tr>
<td>NDA</td>
<td>New drug applications</td>
</tr>
<tr>
<td>NME</td>
<td>New molecular entity</td>
</tr>
<tr>
<td>NSCLC</td>
<td>Non-small cell lung cancer</td>
</tr>
<tr>
<td>Acronym</td>
<td>Definition</td>
</tr>
<tr>
<td>---------</td>
<td>------------------------------------------------</td>
</tr>
<tr>
<td>OAT</td>
<td>Organic anion transporter</td>
</tr>
<tr>
<td>OATP</td>
<td>Organic anion-transporting polypeptide</td>
</tr>
<tr>
<td>OCT</td>
<td>Organic cation transporter</td>
</tr>
<tr>
<td>PBPK</td>
<td>Physiologically based pharmacokinetic</td>
</tr>
<tr>
<td>P-gp</td>
<td>P-glycoprotein</td>
</tr>
<tr>
<td>PhRMA</td>
<td>Pharmaceutical research and manufacturers of America</td>
</tr>
<tr>
<td>Pi</td>
<td>Inorganic phosphate</td>
</tr>
<tr>
<td>PI</td>
<td>Propidium iodide</td>
</tr>
<tr>
<td>P-PBPK</td>
<td>Pediatric PBPK</td>
</tr>
<tr>
<td>PXR</td>
<td>Pregnane X receptor</td>
</tr>
<tr>
<td>QRT-PCR</td>
<td>Quantitative real time polymerase chain reaction</td>
</tr>
<tr>
<td>SLC</td>
<td>Solute carrier</td>
</tr>
<tr>
<td>SP</td>
<td>Side population</td>
</tr>
<tr>
<td>SULT</td>
<td>Sulphotransferase</td>
</tr>
<tr>
<td>TMD</td>
<td>Transmembrane domain</td>
</tr>
<tr>
<td>UGT</td>
<td>Glucuronosyltransferases</td>
</tr>
<tr>
<td>17-AAG</td>
<td>17-(allylamino)-17-demethoxygeldanamycin</td>
</tr>
</tbody>
</table>
CHAPTER 1. INTRODUCTION

1.1. Introduction of Nutlin-3a

1.1.1. Restoring p53 as a therapeutic strategy

Studies from the past three decades show tumor suppressor protein p53 plays important roles in organizing cell defense against cancerous transformation. In responses to stress conditions such as irradiation, DNA damages, and hypoxia, p53 works as a potent transcription factor that activates downstream genes, leading to cell cycle arrest, apoptosis, and inhibition of angiogenesis [1]. p53 can also exert a pro-apoptotic function independent of transcriptional functions [2, 3].

Tumor suppressor p53 is mutated in half of the human tumors [4]. In those tumors that retain wild-type p53, p53 is tightly regulated and maintained at low or undetectable levels. p53 protein has a short half-life of ~20 minutes [5] due to the key negative regulator oncogene murine double minute-2 (MDM2). MDM2 (also known as HDM2) acts as an E3 ubiquitin ligase that facilitates the export of p53 from the nucleus to the cytoplasm and targets p53 for ubiquitin-dependent proteasome degradation [6, 7]. In addition, MDM2 inhibits p53 function by direct binding to the transcriptional binding site of p53, thereby preventing its interaction with the transcription machinery [8]. p53 and MDM2 interact to form an auto-regulatory loop, where increased p53 transcriptionally activates MDM2 and the latter in turn decreases the level of p53 [9] (Figure 1-1).

MDMX (also known as a MDM4 or HDMX), sharing substantial structural homology with MDM2, also has an important role in regulating p53 [10-12]. In addition to inhibiting the transcriptional activity of p53, MDMX forms a heterocomplex with MDM2 that potentiates the ubiquitylation and degradation of p53 [13, 14]. Unlike MDM2, MDMX is not a transcriptional target of p53. Binding of MDMX with MDM2 can cause ubiquitination and degradation of MDMX.

Considering the important roles of MDM2/MDMX in p53 stability and function, restoration of the impaired function of p53 by inhibiting MDM2/MDMX was considered an attractive strategy to treat tumors with wild-type p53 [15]. Several inhibitors of MDM2/MDMX have been discovered and are currently under investigations. For example, a phase I study for JNJ-26854165 (developed by Johnson & Johnson, USA) in patients with advanced stage or refractory solid tumors (ClinicalTrials.gov identifier number: NCT00676910) was completed last year. Phase I clinical trials for ROS45337 (RG7112, developed by F. Hoffmann-La Roche, USA) are ongoing for patients with solid tumors (ClinicalTrials.gov identifier numbers: NCT00659533 and NCT01164033), hematologic neoplasms (ClinicalTrials.gov identifier number: NCT00623870), and liposarcomas (ClinicalTrials.gov identifier number: NCT01143740). Other compounds such as MI-219 [16] (also known as AT219), nutlin-3a [17], SJ-172550 [18], and benzodiazepines [19] are in preclinical development stage.
Figure 1-1. p53 pathway and MDM2/MDMX p53 interaction
1.1.2. Nutlin-3a mechanism of action

In 2004, Vassilev and colleagues reported a group of imidazoline compounds called nutlins are able to inhibit MDM2-p53 binding with high binding potency and selectivity [17]. Nutlin-3 (Figure 1-2) is the most potent compound among the three nutlins (nutlin-1, nutlin-2, and nutlin-3). So far, nutlin-3 is the most widely published small molecule inhibitor of MDM2/MDMX-p53 interaction. Nutlin-3 is a racemic mixture of nutlin-3a (active enantiomer) and nutlin-3b (inactive enantiomer). The binding affinity for nutlin-3a to MDM2 is 150-fold higher than nutlin-3b [17].

Successful development of nutlin is based on understanding the structural biology of the p53-MDM2 interaction. Kussie and colleagues reported a relative deep p53 binding pocket on the surface of the MDM2 protein [20]. Specifically, they found that only three amino acid residues (Phe19, Trp23, and Leu26) of p53 are critical to the binding and fit tightly in the MDM2 binding pocket. This finding made de novo synthesis of small molecule inhibitors of the MDM2-p53 interaction possible. Nutlins were generated by combing structure-based screening of the 3D database, high-throughput screening of large chemical libraries, and extensive chemical modifications of the lead compounds. Crystal structure data of MDM2-nutlin complex proved the binding of nutlin to the p53 pocket [17, 21]. The ethoxy group on the nutlin occupies the position of Phe19, the bromophenyl group occupies the position of Trp23, and the others occupy the position of Leu26 [22]. Since MDM2 and MDMX share structure/sequence similarity and MDMX binds to the similar region of p53 [23], nutlin-3 also binds to MDMX with lower affinity (Figure 1-3) [17, 24].

1.1.3. Reactivation of the p53 pathway by nutlin-3a in vitro

Since 2004, many in vitro studies have been conducted to examine the effect of nutlin-3 on cell cycle arrest and apoptosis. For example, the effect of nutlin-3a and nutlin-3b (1.25-10 μM) on cell cycle arrest was examined in a panel of cancer cell lines from different tumor types, including colorectal (HCT116 and RKO), lung (H460 and A549), breast (MCF7), prostate (LnCaP and 22Rv1), melanoma (LOX), osteosarcoma (SJSA-1), and renal cancer (A498) [25]. 24 hour treatment of nutlin-3a induced a reduction/depletion of the S-phase fraction, as well as G1 and G2 arrest in all the p53 wild-type cell lines tested. Expression of p21, an essential element of p53-induced cell cycle arrest, increased after nutlin-3a treatment. In contrast, these effects were not observed in inactive enantiomer nutlin-3b treatment groups. Colon cancer cells with mutant p53 (HT29) did not respond to nutlin-3a treatment in vitro and in vivo [25].

In contrast to cell cycle arrest, the pro-apoptotic effect is more variable. Apoptosis after nutlin-3a (or -3b) treatment (24~72 hours) was evaluated by Annexin V assay [25]. Annexin V positive fractions varied among p53 wild-type cells from as high as 80% (SJSA-1) to 10% (A549 and HCT116). Since incubation cells with doxorubicin (250 nM) for 48 h led to a dramatic increase of the Annexin V-positive cell fraction in all of the tested lines (including the cell line that had low Annexin V-positive faction after nutlin-3
Figure 1-2.  Chemical structure of nutlin-3

Figure 1-3.  Nutlin-3a binds to the MDM2-p53 and MDMX-p53 binding pockets

treatment), the authors concluded that the low apoptotic level observed after nutlin-3 treatment is not caused by defects in the general components of the apoptotic machinery; rather these cells might have defects in p53-dependent apoptotic signaling.

Studies suggested correlations between high MDM2 expression and strong apoptosis response when p53 wild-type cells are treated with nutlin-3. A significant correlation between MDM2 expression levels and sensitivity to nutlin-3 in p53 wild-type cells was observed in 18 ALL cell lines and 30 primary leukemia samples [26]. Nutlin-3 potently killed wild-type p53 ALL cells over-expressing MDM2. Osteosarcoma cells SJSA-1 and MHM, p53 wild-type cells with 25- and 10- fold MDM2 gene amplification and high MDM2 expression, had the strongest apoptosis response among a panel of 10 p53 wild-type cell lines tested by Annexin-V and microarray analysis [25]. LNCaP (prostate cancer), 22Rv1 (prostate cancer), and RKO (colon cancer) cells with a single copy of MDM2 gene had intermediate levels of apoptotic response. HCT-116 (colon cancer) and U2OS (osteosarcoma) cell lines, lacking the MDM2 gene amplification, had the lowest apoptosis response. Thus, MDM2 expression in tumors may be a valuable response biomarker in the clinic. However, studies for MDM2 might not directly translate to MDMX. In fact, Hu and colleges reported that MDMX over-expression prevents p53 activation by nutlin-3 [27].

In addition, some other characteristics of nutlin-3a are worth mentioning:
1).Unlike radiation and traditional chemotherapy drugs, nutlin-3 activates p53 in a nongenotoxic manner. 2). Nutlin-3 induces apoptosis in p53 wild-type cancer cells; however, it only causes cell cycle arrest in normal cells, which may help protect normal cells from cytotoxic chemotherapies. Thus nutlin-3 was proposed to act as a chemo-protective agent [28]. 3). Multiple studies have suggested synergistic/sensitizing effects of nutlin-3 with radiation [29] or other chemotherapeutic drugs including (but not limited to) topotecan (in retinoblastoma cells) [24], doxorubicin and selumetinib (in acute myeloid leukemia cells) [30, 31], chlorambucil, doxorubicin, fludarabine, dasatinib (in chronic lymphocytic leukemia cells) [32-35], R-roscovitine (in neuroblastoma cells) [36], vincristine, actinomycin D, doxorubicin, etoposide (in Ewing sarcoma cells), and bortezomib (in myeloma, thyroid, breast, and prostate carcinomas and colon carcinoma cells) [37, 38].

1.1.4. **In vivo anti-tumor effect of nutlin-3a**

*In vivo*, nutlin-3a monotherapy demonstrated anti-tumor efficacy in preclinical models of human osteosarcoma, prostate cancer, retinoblastoma, KSHV lymphoma, and neuroblastoma with wild-type p53 [17, 24, 25, 39-41]. Vassilev et al. first reported the *in vivo* activity of nutlin-3 in nude mice bearing subcutaneous human osteosarcoma xenograft (SJSA-1). Nutlin-3 (po. 200mg/kg BID for 3 weeks) was well tolerated without causing significant weight loss or any gross abnormalities upon necropsy at the end of the treatment. Compared to the vehicle control group, nutlin-3 treatment resulted in 90% inhibition of tumor growth [17].
Tovar et al. conducted *in vivo* study of nutlin-3a in nude mice bearing SJSA-1 (osteosarcoma), MHM (osteosarcoma), LNCaP (prostate cancer), 22Rv1 (prostate cancer) and HT29 (colon cancer) tumors [25]. SJSA-1-bearing mice were treated with an oral dose of 50mg/kg, 100mg/kg or 200mg/kg nutlin-3a twice daily for 3 weeks. Nutlin-3a dose dependently suppressed SJSA-1 tumor growth, with substantial tumor shrinkage observed in the 200mg/kg treatment group. The 200 mg/kg oral nutlin-3a twice daily regimen was also efficacious in MHM (3 weeks treatment), LNCaP (2 weeks treatment), and 22Rv1 (2 weeks treatment) models with average tumor growth inhibition > 98%. In p53 mutant HT29 xenograft, nutlin-3a did not reduce the tumor size. The data showed a reasonable correlation between *in vitro* and *in vivo* tumor response. Similar to the report from Vassilev et al., no weight loss or significant pathological changes were observed during the study.

Laurie et al. conducted the first *in vivo* study to assess the effect of nutlin-3 on retinoblastoma [24]. Subconjunctival injections of 1 μl nutlin-3 (170 mM) and topotecan (2 mM) both as a single agent and in combination were administered into each eye of tumor-bearing mice daily for 5 days. Total treatment amount per eye was 85 pmol nutlin-3 and 2 nmol topotecan. Both nutlin-3 and topotecan were effective as a single agent in the Y79-luc orthotopic model. The combination of subconjunctival topotecan and nutlin-3 resulted in an 82-fold tumor burden reduction with no ocular or systemic side-effects. Brennan et al. recently reported a study aimed to identify better chemotherapeutic combinations for the treatment of retinoblastoma in genetically engineered mouse models and orthotopic xenograft models of human retinoblastoma [42]. SCID mice bearing SJ-39 retinoblastoma tumor cells received vincristine/toposide/carboplatin, carboplatin(subconj)/topotecan(syst), or carboplatin(subconj)/topotecan(syst) alternating with nutlin-3a(OC)/topotecan(syst). The nutlin-3a(OC)/topotecan(syst)-containing group showed significantly better response. Subconjunctival administrations of nutlin-3a alone or in combination with topotecan were well tolerated without ocular or systemic toxicity.

Van Maerken et al. reported the effect of nutlin-3 on nude mice bearing chemo-resistant, MYCN-amplified neuroblastoma [43]. 200mg/kg oral nutlin-3 twice daily treatment reduced tumor growth and metastasis in the p53 wild-type UKF-NB-3rDOX20 xenograft without causing signs of toxicity. No treatment effect was observed in p53 mutant UKF-NB-3rVCR10 xenograft, suggesting p53 status significantly influences the *in vivo* response to nutlin-3 treatment.

### 1.2. ABC Transporters and Drug Interaction

#### 1.2.1. ABC transporter family

Transporters are membrane proteins that play important roles in controlling the influx and efflux of ions, glucose, bile acids, vitamins, hormones, lipids, fatty acids, toxins, and drugs across cell membranes [44, 45]. 5~7% (> 2,000) of all human genes
code for transporters or transporter-related proteins [46]. Among all the transporters, ATP-binding cassette (ABC) and Solute carrier (SLC) transporters are two major families of membrane proteins that are important for transporting drugs. So far, more than 400 membrane transporters in these two families have been annotated in the human genome [47].

ABC transporters are a family of active transporters relying on adenosine triphosphate (ATP) hydrolysis to pump substrates in (influx) and out (efflux) of the cell membranes. In prokaryotes, ABC transporters function as both uptake transporters and efflux transporters. However, in eukaryotes, ABC transporters function only as efflux transporters [48, 49]. The ABC gene family is composed of 49 genes in 8 subfamilies in the human genome [45]. The basic structure of ABC transporters contains two types of domains: nucleotide binding domain (NBD) and transmembrane domain (TMD) (Figure 1-4) [50]. The NBD, the conserved domain among the ABC transporters, plays a critical role in ATP binding and hydrolysis. Unlike NBD, TMD varies significantly in terms of the sequence, length, and number of transmembrane helices [51]. TMD binds to substrates and determines the transporter specificity through substrate-binding sites [52]. Among the ABC family of transporters, P-glycoprotein (P-gp/MDR1/ABCB1), multidrug resistance protein 1 (MRP1/ABCC1), and breast cancer resistance protein (BCRP/MXR/ABCG2) are three major members associated with multidrug resistance [53].

P-gp, the product of MDR1 (or ABCB1) gene, was one of the first members of the ABC superfamily studied. Long before P-gp was discovered, it had been reported that incubating cancer cells with chemotherapy agents will generate subline cells that are resistant to not only the selecting agents but also to other structurally different agents [53-55]. 10 years after the first report of the 170 kDa glycoprotein [53-55], Roninson et al. reported the cloning of the gene encoding P-gp [56]. P-gp contains 2 NBDs and 12 TMDs (Figure 1-4) [50]. In addition to tumor cells, P-gp is also expressed in multiple normal organs/cells, such as intestinal enterocytes, kidney proximal tubule, hepatocytes (canalicular), and brain capillary endothelial cells. P-gp transports a broad variety of substrates out of the cells, including endogenous substrates, and drugs such as vincristine, vinblastine, doxorubicin, topotecan, mitoxantrone, etoposide, paclitaxel, docetaxel, and digoxin.

BCRP, a 72-kDa protein product of ABCG2 gene, is also called ABCP or MXR. As indicated by its most commonly used name, BCRP was identified from a multidrug-resistant human breast cancer subline (MCF-7/AdrVp) in 1998. In 1999, it was cloned from mitoxantrone selected cells; thus it was also named mitoxantrone resistance protein (MXR) [57]. Interestingly, the clones from the drug selected cells containing single nucleotide mutations at the position of amino acid 482 (R for wild-type protein, T in BCRP, and G in MXR), causing changes in substrate specificity [50, 58]. Unlike P-gp and MRPs, BCRP is a “half-transporter.” It contains only 1 NBD and 6 TMDs (Figure 1-4) [50]. BCRP is expressed in tumor cells, hematopoietic stem cells, placenta, small intestine, mammary glands, testis, liver, blood brain barrier, and the adrenal gland.
Figure 1-4. The structures of three categories of ABC transporter

a. ABC transporters such as multidrug resistance MDR1 and multidrug resistance associated protein 4 (MRP4) have 12 transmembrane domains and two ATP binding sites.

b. The structures of MRP1, 2, 3 and 6 are similar in that they possess two ATP binding regions. They also contain an additional domain that is composed of five transmembrane segments at the amino-terminal end, giving them a total of 17 transmembrane domains.

c. The ‘half-transporter’ ABCG2 contains 6 transmembrane domains and one ATP-binding region — in this case, on the amino-terminal side (N) of the transmembrane domain. In other ‘half-transporters’, such as the transporter associated with antigen processing, the ATP-binding cassette is found on the carboxy-terminal (C) side of the transmembrane domain. Half-transporters are thought to homodimerize or heterodimerize to function.

BCRP transports endogenous substrates and drugs such as mitoxantrone, topotecan, SN-38, methotrexate, doxorubicin, and daunorubicin [50, 58].

In addition, P-gp, BCRP, and many other members of ABC transporter family play important roles in absorption, distribution, metabolism, and excretion (ADME) of chemotherapy drugs that are substrates of the ABC transporters.

1.2.2. Role of ABC transporter in drug ADME

At least 10 ABC transporters (P-gp, MDR3, BSEP, MRP1, MRP2, MRP3, MRP4, MRP5, MRP6, and BCRP) are involved in drug disposition [47, 59]. They are located on the apical or basolateral side of endothelial or epithelial cells in various organs (Table 1-1) [47, 59].

1.2.2.1. Effects on drug absorption. ABC transporters located on the apical membrane of enterocytes, such as P-gp, MRP2, and BCRP [47], can pump their substrates back to the intestinal lumen—thus limiting the absorption of some orally administered drugs. For example, paclitaxel bioavailability in P-gp knockout mice increased from 11.2% to 35.2% compared to the wild-type mice [60]. Docetaxel bioavailability increased from 3.6% to 22.7% in P-gp knockout mice compared to the wild-type mice [61]. Leggas et al. reported that topotecan bioavailability increased in both BCRP and P-gp knockout mice and further increased with gefitinib (an inhibitor of BCRP and P-gp) treatment [62]. MRP1 and MRP3 are expressed at the basolateral side. Thus, theoretically, they could increase absorption of some drugs. A recent study suggests that MRP1 might facilitate the absorption of cobalamin in mice [63]. Several ABC transporters are expressed in the lungs and therefore may potentially affect the absorption of inhaled drugs. However, in vivo evidence is needed to support this hypothesis [64].

1.2.2.2. Effects on drug distribution. ABC transporters located on the blood-brain barrier (BBB), blood-cerebrospinal fluid (CSF) barrier, blood-placental barrier, blood-testis barrier [65], blood-retina barrier [66], and heart [67] can limit the drug distribution to corresponding organs. BBB is a good example on how ABC transporters affect drug distribution. Before the 1990s, the BBB was considered a physical barrier formed by tight junctions between brain capillary endothelial cells (BCEC) that lack fenestrations [68]. Now, it is well established that P-gp on the apical membrane of BCECs is also an important component of BBB. The brain accumulation of many P-gp substrates can be much higher in P-gp knock-out mice than in wild-type mice, and inhibiting P-gp can increase the brain distribution. For example, amprenavir, an HIV protease inhibitor, is a substrate of P-gp [69] but not a substrate of BCRP [70]. Brain concentrations of [14C]-amprenavir were 27-fold higher in mdr1a-/-/1b-/- mice compared to the wild-type mice. In the presence of P-gp and BCRP inhibitor GF120918, a 13-fold increase of [14C]-amprenavir brain concentrations compared to the vehicle control treated
Table 1-1. Important ABC transporters

<table>
<thead>
<tr>
<th>Name</th>
<th>Polarity</th>
<th>Locations</th>
</tr>
</thead>
<tbody>
<tr>
<td>P-gp</td>
<td>Apical</td>
<td>Kidney, adrenal gland, liver, pancreas, intestine, lung, BBB, placenta, prostate, skin, heart, skeletal muscle, ovary, testis, retina</td>
</tr>
<tr>
<td>MDR3</td>
<td>Apical</td>
<td>Liver</td>
</tr>
<tr>
<td>BCRP</td>
<td>Apical</td>
<td>Placenta, mammary gland, BBB, liver, intestine, kidney, lung</td>
</tr>
<tr>
<td>MRP1</td>
<td>Apical (placenta, BBB) Basolateral (others)</td>
<td>Kidney, lung, testis, skeletal muscle, heart, placenta, liver, intestine, brain</td>
</tr>
<tr>
<td>MRP2</td>
<td>Apical</td>
<td>Liver, kidney, intestine, placenta</td>
</tr>
<tr>
<td>MRP3</td>
<td>Basolateral</td>
<td>Adrenal gland, intestine, pancreas, intestine, gallbladder, placenta, liver, kidney</td>
</tr>
<tr>
<td>MRP4</td>
<td>Apical (kidney, BBB) Basolateral (prostate, choroid plexus)</td>
<td>Prostate, kidney, liver, brain, pancreas</td>
</tr>
<tr>
<td>MRP5</td>
<td>Apical (BBB) Basolateral (others)</td>
<td>Heart, brain, neurons</td>
</tr>
<tr>
<td>MRP6</td>
<td>Basolateral</td>
<td>Liver, kidney, skin, lung, heart, intestine, pancreas, stomach</td>
</tr>
<tr>
<td>BSEP</td>
<td>Apical</td>
<td>Liver</td>
</tr>
</tbody>
</table>

Sources: Adapted with permission.


group was observed [71]. Blood concentrations of $[^{14}\text{C}]-$amprenavir increased 1.1-fold and 2-fold in $m\text{d}r1a^{-/-}/1b^{-/-}$ mice and GF120918 treated mice, respectively [71]. Although less prominent than P-gp, BCRP and some MRPs are also involved in the function of BBB. These transporters can protect the brain from peripheral toxins but also hinder the delivery of central nervous system (CNS) drugs.

1.2.2.3. Effects on drug metabolism. ABC transporters do not metabolize drugs themselves, but they affect metabolic clearance remarkably through interplay with drug metabolism enzymes. It has been proposed that ABC transporters and drug metabolism enzymes have undergone co-evolution toward a united xenobiotic defense system [45]. The relationship between P-gp and CYP3A4 is considered evidence of this theory. The $ABCBI$ gene (encodes for P-gp) and the cluster of $CYP3A4$ genes are both at chromosome 7 and just 119kb apart [45]. In addition, both genes are regulated by pregnane X receptor (PXR). P-gp and CYP3A4 also share similar substrate specificity and are co-localized in important drug-eliminating organs such as liver, kidney, intestine, and lung [45, 72]. The most well-known effect of P-gp on CYP3A4 is exemplified by the intestinal first-pass metabolism. As mentioned, P-gp in enterocytes can pump back its substrate into the gut lumen, thus decreasing the fraction of absorption (fa). In addition, the P-gp-mediated efflux also helps decrease the likelihood of CYP3A4 saturation by lowering the intracellular peak drug concentration. The effluxed drug may undergo re-absorption, but the overall effect will be more opportunities for the CYP3A4 to metabolize the substrate drug. Therefore, P-gp can indirectly decrease the fraction of the intact drug escaping gut metabolism (fg) [59, 68, 73]. Drug conjugates formed by Phase II drug metabolism enzymes, e.g., UDP-glucuronosyltransferases (UGTs), sulfotransferases (SULTs), and glutathione S-transferases (GSTs), can also be effluxed by MRP2 and BCRP. This process is sometimes called “Phase III drug metabolism”.

1.2.2.4. Effects on drug excretion. Drug excretion refers to the final removal of intact drugs or their metabolites from the body. Although this step can take place in several organs, biliary excretion and renal excretion are the most important routes [59]. Different ABC transporters are localized at the canicular or sinusoidal membrane of hepatocytes and in the kidney. Sinusoidal membrane transporters MRP1, MRP3, MRP4, and MRP6 extrude some drug metabolites (in most cases) or intact drugs back to the blood, and present them to bile or renal excretion. On the other hand, canicular membrane transporters P-gp, MRP2, and BCRP excrete their substrates directly into bile. P-gp is mainly responsible for cationic drugs or metabolites, while MRP2 and BCRP are for anionic drugs or metabolites [59]. However, there are also some controversies on the importance of MRP2 and BCRP, since it is difficult to differentiate their contribution from OATPs. ABC transporters are also located at both the apical and basolateral membrane of the renal epithelial cells. P-gp, MRP2, and MRP4 are expressed at the proximal tubular basolateral membrane facilitating the excretion of compounds into the urine. It seems that P-gp is responsible for excretion of digoxin and some hydrophobic cationic drugs, while MRP2 and MRP4 are for anionic drugs or metabolites. MRP1 is expressed at the apical cell membrane of distal tubules and collecting ducts. It may be
part of a mechanism to prevent drug accumulation (which is toxic to nephron) after water re-absorption in that area.

### 1.2.3. Role of ABC transporter on drug resistance

Resistance to chemotherapy is one of the major hindrances to the current multimodal cancer treatment paradigms. The resistance can be either a result of changes in drug’s ADME at a non-cellular level or the consequence of certain mechanisms within tumor cancer cells [74]. As implied by the names of their well-known members (e.g., Breast Cancer Resistance Protein, Multidrug Resistance Proteins, as well as the gene name of P-gp, Multidrug Resistance gene 1), ABC transporters are involved in drug resistance in cancers.

As mentioned in the previous section, some ABC transporters affect ADME of chemotherapeutic agents. Therefore, induction of these transporters by the chemotherapeutic agent itself or other co-administrated drugs can cause resistance to that agent at the non-cellular level. This can work together with other drug resistance mechanisms at this level, e.g., drug metabolism enzymes induction in drug-handling organs as well as poor vascularization and acidic pH in solid tumors [75, 76].

Cellular drug resistance can be further stratified into two types: the resistance to a class of drugs with a similar mechanism of action, and the resistance to various drugs with different structures and targets [75]. The former type of resistance obviously results from changes in drug targets, while the latter type, so-called multiple drug resistance (MDR), has been a topic of much discussion.

Multiple mechanisms have been proposed for MDR, including reduced sensitivity to apoptosis and existence of cancer stem cells, as well as high levels of expressions of drug metabolism enzymes (such as GST) and ABC transporters in cancer cells [74, 76]. MDR caused by ABC transporters is called “transport MDR” or “classical MDR” while MDR by other mechanisms is referred as “atypical MDR” or “non-classical MDR” [74, 76].

All 10 aforementioned ABC transporters that are important for ADME plus 3 additional members (MRP7, MRP8, and ABCA2) have been shown to cause drug resistance in cell lines *in vitro* [77, 78]. In many different cancer tissues from both drug-naïve and treated patients, various ABC transporters frequently can be detected [79]. However, it is more challenging to get a clear picture of their significance in clinical MDR. In order to discern the contribution of ABC transporters from other factors, correlation analysis between poor chemotherapy outcome and expression of ABC transporters in a large cohort of cancer patients is a necessity. Another reason that makes the consensus on clinical relevance of some transporters hard to achieve is the large variety of tumor types and drug treatments [79].
Currently, most studies on the relationship between ABC transporters and clinical MDR are focused on P-gp, MRP1, and BCRP, which are also the most extensively studied ones in drug ADME. Correlation between P-gp expression and prognosis of chemotherapy in breast cancer, sarcoma, and acute myelogenous leukemia (AML) are well established [80]. However, the results of clinical trials using specific P-gp inhibitors are not very promising [81].

The expression level of MRP1 does not correlate with the clinical MDR of AML. Its significance on the chemotherapy outcome of chronic lymphocytic and promyelocytic leukemia, non-small cell lung cancer (NSCLC), and breast cancer has been a point of controversy [80]. There are conflicting data regarding the role of BCRP in clinical MDR in leukemia [53, 80, 81]. No or little clinical relevance of BCRP expression was found in breast cancer, ovarian cancer, and locally advanced bladder cancer. Correlation between BCRP expression and adverse prognosis of lung cancer, esophageal cancers, and some lymphomas has been reported [82].

Recently, the cancer stem cells paradigm has been incorporated into the drug resistance concept. Cancer stem cells are drug-resistant pluripotent cells expressing high levels of ABC transporters, especially BCRP. This subpopulation can survive chemotherapy when the other committed non-resistant cancer cells in the original tumor mass are killed and therefore serve as an unrestricted reservoir for drug resistant tumor relapse [80, 83]. Thus, some conclusions on the relevance of BCRP in clinical MDR need to be re-evaluated, since BCRP expression in cancer stem cells was not examined in many previous solid tumor studies [82]. Some researchers have proposed that the failure in P-gp inhibitor development may be due to missing an important target—the BCRP [84].

1.2.4. Mechanism of action of ABC transporters

Current understanding of the mechanism of action of ABC transporters is briefly summarized below (Figure 1-5) [50, 51, 85-90]. As mentioned before, TMDs of ABC transporters bind to drug substrates, while NBDs bind to ATP. Binding of substrate to the TMD stimulates the ATPase activity. ATPase facilitates ATP hydrolysis and releasing of inorganic phosphate (Pi) and adenosine diphosphate (ADP). ATP hydrolysis provides the energy to cause a conformational change at the TMDs. TMDs will change from inward-facing conformation (facing the inside) to outward-facing conformation (facing the outside) and release substrate to the extracellular space. Experimental data supported that two ATP hydrolysis events are needed to transport one drug molecule. These two events do not happen simultaneously. The first ATP hydrolysis is needed to transport the substrate and the second ATP hydrolysis is needed to “reset” the transporter from outward facing back to forward facing so that the transporter can bind substrate again [91]. Formation of a homodimer or heterodimer is a prerequisite for functionality of half-transporters such as BCRP [50, 58, 92].

Based on this mechanism of action, the ATPase assay was developed to indirectly
Figure 1-5.  Mechanism of action of a typical ABC transporter

ATP dependent closure/dimerization of cytosolic NBDs provides the power stroke that pulls the TMDs from an inward- to outward facing conformation. Both exporters and importers probably use the same basic mechanism but shift which state binds the transport substrate (red) with high affinity. Most eukaryotic ABC transporters are heterodimers, with 2 homologous but nonequivalent halves (green and blue).

detect the activity of ABC transporters to determine substrates and inhibitors of ABC transporters. The ATPase assay contains two modes, activation mode and inhibition mode. Under activation mode, ABC transporter substrates bind to the TMD and stimulate ATPase activity, which further hydrolyzes ATP and releases Pi, which can be detected by a colorimetric reaction and is proportional to the ATPase activity. Increased Pi release suggests that a compound is a substrate of ABC transporters. Under inhibition mode, Pi release is detected in the presence of a known ABC transporter substrate with or without a test compound. If the test compound is an inhibitor of the transporter or slowly transported substrate, Pi release stimulated by the transporter substrate will be reduced [93, 94].

1.2.5. Effect of p53 on ABC transporters

Conflicting data suggested that p53 may also regulate the expression of ABC transporters [95, 96]. In 1992, Chin et al. first reported that the MDR1 promoter was repressed by wild-type p53 [97]. Later, Stauss et al. reported that the MDR1 downstream promoter contains a wild-type p53-binding site [98]. Unlike wild-type p53, mutant p53 was found to activate the MDR1 promoter [97]. Similarly, Zastawny et al. also reported that wild-type p53 represses MDR1 promoter activity, and mutant p53 enhances MDR1 promoter activity [99]. Further study showed the overlapping region on the MDR1 promoter for mutant p53 transactivation and for basal promoter activity [100]. A study by Sampath et al. suggested while mutant p53 requires an Ets-1 transcriptional factor binding site to regulate MDR1 promoter transcriptionally, wild-type p53 does not interact with Ets-1 [101]. The increased MDR1 expression could be caused by both a loss of p53 repression on MDR1 promoter and an increased transactivation of MDR1 through a “gain of function” of mutant p53 [102]. However, the role of p53 in regulating MDR1 is still controversial [95, 102]. Transfection of wild-type p53 expression had been reported to stimulate MDR1 promoter in p53 negative cell lines [103] or show no (or marginal) change in MDR1 gene expression and function [104, 105].

In addition to MDR1, wild-type p53 has been reported to suppress MRP1 reporter activity [106]. Unlike MDR1, mutant p53 did not up-regulate MRP1 [101, 106]. Data about regulation of MRP1 by p53 is also conflicting. Wild-type p53 enhanced MRP expression and activity instead of suppressing MRP1 expression in three of the five cell lines, and no change was observed in other two of the five cell lines [96]. Recently, Wang et al. reported that wild-type, but not mutant p53, can reduce the expression of BCRP in MCF-7 cells [107].

Regulation of ABC transporter expression by p53 can have a functional effect on drug resistance. For example, wild-type p53 inactivation (introduced by expressing dominant negative mutant p53) increased the expression of MDR1. This leads to decreased uptake and increased resistance to vinblastine [102]. Increased vincristine sensitivity after changing the cells from expressing dominate negative mutant p53 to wild-type p53 was also reported [96].
1.2.6. Evaluation of transporter mediated drug-drug interaction

Because of the important roles of transporters on drug disposition, inhibition or activation of ABC transporter(s) may alter the PK of a compound that is substrate(s) of ABC transporter(s). For example, oral co-administration of P-gp inhibitor clarithromycin can lead to a 64% increase of AUC of P-gp substrate digoxin, which explains the clinical cases of clarithromycin-induced digoxin toxicity [108]. P-gp inhibitor dronedarone resulted in a 157% increase of digoxin AUC [47]. BCRP and P-gp inhibitor GF120918 resulted in a 143% increase of oral topotecan AUC [47]. Therefore, in vitro and sometimes in vivo follow-up studies are needed to understand transporter-mediated drug-drug interaction for avoiding serious adverse events.

In 2006, the Food and Drug Administration (FDA) published guidance for studying transporter-based drug-drug interactions. A decision tree regarding P-gp-mediated drug-drug interaction was included in the guidance [47]. In 2010, the International Transporter Consortium further broadened the decision tree (Figure 1-6) [47]. Approaches to answer important questions were addressed for both ABC transporters (P-gp and BCRP) and SLC transporters (organic cation transporter (OCT) and the organic anion transporter (OAT)) [47]. These questions include, for example, which transporters are important for drug absorption and disposition, what in vitro methods are recommended to evaluate drug-transporter interaction, and what are the criteria for conducting clinical studies based on in vitro data.

For P-gp and BCRP, the bidirectional transport assay using polarized monolayer cells is one of the important preferred functional studies to identify if a compound is a substrate and/or inhibitor of P-gp. Caco-2 cells are derived from human epithelial colorectal adenocarcinoma cells. When cultured under specific conditions, Caco-2 cells become differentiated, polarized, and functionally resemble the morphology, polarity, and expression patterns of the transporters and enzymes of the small intestine [109]. Caco-2 cells have been widely used as an in vitro model to evaluate intestinal drug absorption and efflux. Transporter transfected Madin-Darby canine kidney epithelial (MDCK) cells or porcine kidney epithelial cells (LLCPK1) cells are also routinely used in bidirectional transport assay. Similar to Caco-2 cells, these cells can also polarize and form a monolayer. Because P-gp and BCRP are expressed at the apical side of the polarized cells [109], B to A transport represents the passive diffusion and the efflux of a compound, and A to B transport represents the passive diffusion and the uptake of a compound. Therefore, it is important to look at both A to B and B to A direction for evaluating P-gp and BCRP efflux in the system. When interpreting data from these systems, one must consider that multiple endogenous drug transporters are expressed in Caco-2 (P-gp, BCRP, MRP1-6, ABCA1, ABCG1, HPT1 and many SLC family uptake transporters) [110], MDCK II (P-gp, MRPI, MRP2, and MRP5) [111, 112], and LLC-PK1 (P-gp, MRPI and MRP2) [113].

A decision tree for P-gp or BCRP substrate interactions is listed in Figure 1-6 [47]. An efflux ratio less than 2 indicates poor or non-P-gp (or non-BCRP) substrate. For
A. Decision tree for P-gp or BCRP substrate interaction. A new molecular entity (NME) is considered to be a potential P-gp or BCRP substrate if the efflux ratio — basal to apical (B-A) to apical to basal (A-B) — is ≥ 2 in an epithelia cell system that expresses one or both transporters (see (a) in the figure). A net flux ratio cut-off higher than 2 or a relative ratio to positive controls may be used to avoid false positives if a ratio of 2 is deemed non-discriminative as supported by prior experience with the cell system used. Additional corroboration that an NME may be a P-gp or BCRP substrate can be achieved with the use of inhibitors. Reduction of the flux ratio by the P-gp (or BCRP) inhibitors should be greater than 50% (see (b) in the figure). If the flux ratio is not reduced by P-gp (or BCRP) inhibitors, then other efflux transporters may be responsible for the observed net flux (see (d) in the figure).

B. Decision tree for transporter inhibitor interaction. [I]1 is the steady-state total Cmax at the highest clinical dose, and [I]2 is the theoretical maximal gastrointestinal drug concentration after oral formulation at the highest clinical dose in a volume of 250 ml.

A.

In bi-directional transporter assays, for example, in Caco-2 or P-gp-overexpressing polarized epithelial cell lines, is the net flux ratio of NME ≥ 2?

- **Net flux ratio ≥ 2**
- **Net flux ratio < 2**

b. Is efflux significantly inhibited by 1 or more P-gp inhibitors?

- **Yes**
  - Probably P-gp substrate

- **No**
  - Other efflux transporters are responsible for observed data

Complete an assessment of preclinical and clinical information to determine whether an *in vivo* DDI study is warranted.

B.

Bi-directional transporter assay with a probe P-gp substrate. For example, in Caco-2 or P-gp-overexpressing polarized epithelial cell lines

Net flux ratio of a probe substrate decreases with increased concentrations of the investigational drug

- **Net flux ratio of the probe substrate is not affected with increased concentrations of the investigational drug**
  - Poor or non-inhibitor

Determine $K_i$ or $IC_{50}$ of the inhibitor

- **$[I]/IC_{50}$ (or $K_i$) ≥ 0.1 or $[I]/IC_{50}$ (or $K_i$) ≥ 10**
  - An *in vivo* drug interaction study with a P-gp substrate such as digoxin is recommended

- **$[I]/IC_{50}$ (or $K_i$) < 0.1 and $[I]/IC_{50}$ (or $K_i$) < 10**
  - An *in vivo* drug interaction study with a P-gp substrate may not be needed
compounds with efflux ratio ≥ 2, inhibitor(s) will be used. If the efflux ratio reduction is larger than 50 % or the efflux ratio close to 1 can be achieved after adding P-gp (or BCRP) inhibitor(s) at effective concentration, the compound is probably a substrate of P-gp (or BCRP). Otherwise, other efflux transporters may be responsible for the high efflux ratio. If an in vitro study suggested that a compound is a substrate of P-gp or BCRP, an in vivo drug-drug interaction study may be needed, depending on the property of a compound. For a drug that falls into Biopharmaceutics Classification System (BCS) Class I (high solubility, high permeability) or Biopharmaceutics Drug Disposition Classification System (BDDCS) Class I (high solubility, extensive metabolism), even if it could be P-gp substrate in vitro (for example, verapamil), its bioavailability is not likely to be affected by P-gp modulators [114]. If disposition in kidney, brain, and tumor is not important for that drug, then no in vivo studies on drug-drug interaction are necessary [114].

If a compound reduced the efflux ratio of P-gp (or BCRP) in bi-directional transport assay, this compound might be a potential P-gp (or BCRP) inhibitor in vivo. The International Transporter Consortium recommended \( \frac{I_1}{IC_{50}} \geq 0.1 \) or \( \frac{I_2}{IC_{50}} > 10 \) as cut-off values for further in vivo drug interaction studies (Figure 1-6B) [47]. \( IC_{50} \) is the in vitro IC50, \( I_1 \) is the steady-state unbound Cmax at the highest clinical dose, and \( I_2 \) is the theoretical maximal gastrointestinal drug concentration after oral formulation at the highest clinical dose in a volume of 250 ml. If \( \frac{I_1}{IC_{50}} \geq 0.1 \) or \( \frac{I_2}{IC_{50}} > 10 \) occur, an in vivo drug interaction study with P-gp or BCRP substrate is recommended. One possible substrate for an in vivo P-gp-mediated drug interaction study is digoxin, and possible substrates for an in vivo BCRP-mediated drug interaction study are sulphasalazine, rosvuastatin, pitavastatin, ciproflozacin, and dipyridamole.

1.3. Physiologically-based Pharmacokinetic (PBPK) Modeling and Simulation in Drug Development

1.3.1. Introduction of PBPK model

1.3.1.1. Classical and PBPK model. Disposition of a drug by the body is a complex process. A compound can be administered by various routes such as oral, intravenous, subcutaneous, intramuscular, or subconjunctival injection. Through various mechanisms of absorption, the drug is absorbed and then distributed to the tissues by the body fluids (e.g., blood, lymphatic system, CSF). The drug can be metabolized in the liver and/or other tissues to other compound(s) and/or eliminated unchanged through bile or the kidney. Protein binding in the plasma and different tissues can be different due to various protein contents. As discussed earlier, efflux and uptake transporters expressed throughout the body can impact the ADME process. In the same tissue, drug concentrations in the vascular, extracellular, and intracellular compartments can be different. Due to the complexity of this process, drug concentrations in the blood and in
different parts of the body can be quite variable. Thus drug concentrations in the plasma do not always reflect target tissue concentrations.

An appropriate pharmacokinetic model that adequately describes and reasonably predicts the time-dependent drug concentrations in the body is critical in both preclinical and clinical drug development. Currently, two broad approaches to pharmacokinetic analysis are classical (sometimes called empirical or conventional) and PBPK models, although the two approaches do share similar features.

The classical pharmacokinetic model focuses mainly on the drug concentrations in blood (whole blood, plasma, or serum) or other easily accessible body fluids (urine, feces, CSF, or breast milk). It generalizes the complex drug distribution process into multiple theoretical compartments. An example of classic PK model structure is provided in Figure 1-7. Compartments in classical PK models do not represent real organ compartments and usually lack mechanistic insight. This limits their application in certain areas of drug discovery and development such as predicting human systemic and target tissue drug concentrations based on in vitro data or animal PK data, and predicting drug-drug interactions.

PBPK models, on the other hand, are more complex models that map the drug PK process onto physiologically realistic compartment structures. A typical structure of a PBPK model is given in Figure 1-8 [115]. In such a model, the body is usually modeled as a closed circulatory system consisting of tissues that are important for drug absorption, distribution, metabolism, elimination, or any other tissues that are of interest. Compartments that represent real tissues are connected by blood flow. In general, tissue-specific arterial blood flow serves as model input into the tissues and comes out of the tissues as venous blood, thus serving as model output, with some exceptions. For splanchnic organs such as stomach, intestine, spleen and pancreas, venous blood flows into the liver through the portal system. For the lung, the pooled venous blood flows in the lung and arterial blood flows out of the lung. In contrast to the classic PK model, the physiologically-based nature of the PBPK modeling allows us to address mechanistic questions with regard to the PK as well as extrapolating knowledge obtained from one species to another, including human.

However, the distinction between the classic and PBPK model is not always clear. As Nestorov mentioned in his reviews, it is impossible and unnecessary to define formally what a PBPK model is or to specify a clear distinction between the classic and PBPK model [116, 117]. The classic model (especially mechanistic PK model) contains more and more physiological information (e.g., including body weight in the allometric scaling, using bile empty time for modeling enterohepatic circulation). And the PBPK model usually incorporates classic PK model components. For example, PBPK modeling software PK-sim® incorporates the animal clearance value from classic PK model into the PBPK model to predict concentration time profile both in animal and in human. Also, one important step for establishing the whole body PBPK model is to generate PK model in the plasma using classic PK modeling. To distinguish between a predominantly classical and predominantly PBPK model, the convention is to look at the model
Figure 1-7. Example model structure of the classical PK model
Figure 1-8. Typical structure of a whole-body PBPK model

Q refers to blood flow: to the lungs (Qpul), the heart (Qca), the kidneys (Qre), the bones (Qbo), the muscles (Qmu), the spleen (Qsp), the liver (Qha), the hepatic vein (Qhv), the gut (Qgu), the fat (Qfa), the skin (Qsk), and the thymus (Qth).

structure. As mentioned in a review by Nestorov, if the model structure precedes the analysis of the compound specific data, and predominantly represents actual tissue or organ spaces, then it can be classified as a PBPK model [117]. It should be mentioned that both the classical PK model and the PBPK model belong to the compartment PK model hierarchy because both models classify the body into a number of subunits called “compartments” [117, 118].

1.3.1.2. PBPK application area. Due to the advances in computing power, PBPK models have been increasingly used in recent years [119]. The majority of all PBPK related publications (60%) deal with issues pertaining to risk assessment of environmental chemicals [120]. Because the PBPK model is more mechanistically, anatomically, and physiologically relevant than the classic compartment modeling, the established PBPK model from animal study can be used to extrapolate to humans.

In the drug discovery and development area, there is a particular interest to use the PBPK model for estimating the human PK of drug candidates from in silico, in vitro, or in vivo animal PK data and to select the most promising compounds for further development. Methods for predicting human PK are quite variable, and many methods are under evaluation. For example, detailed scaling methods can be found in recently published papers from the Pharmaceutical Research and Manufacturers of America (PhRMA) group, including 24, 29, and 66 methods for predicting human volume of distribution, human clearance and oral area under the curve, respectively [121-125], based on in vitro and animal PK data. PBPK model has also been used for special population—such as pediatrics [126-134], elderly patients [135, 136], pregnant [137-139] and lactating [140, 141] individuals, and for patients with organ impairments [142]. It has been applied to predict human drug-drug interaction [130, 143-151], inter-individual variability [120], and the effect of genetics [152-154]. In addition to small molecules, the applications of the PBPK model have also been extended to large molecules [155-163] and nano-particles [164-167].

1.3.1.3. PBPK model software. Software is a critical tool for establishing PBPK models and performing simulations. There are a number of commercial software products developed for building PBPK models, including Simcyp (Simcyp Ltd., Sheffield, UK), GastroPlus™ (Simulations Plus Inc, Lancaster, CA) and PK-Sim® (Bayer Technology Services, Leverkusen Germany). These products usually do not require code writing and are capable of performing complex IVIVE simulation, DDI prediction, and extrapolating PK parameters from adults to pediatric population without in vivo experiment data. Software such as SAMM II, NONMEM, ADAPT, and Phoenix (WinNonlin 6) can be used for both conventional PK and PBPK modeling. Different software requires different coding. In some cases, such as in SAAMII and Phoenix WinNonlin 6, writing code may not be needed. Software products such as MATLAB have powerful programming flexibilities.
1.3.2. Basic concepts of PBPK modeling

1.3.2.1. Major components of PBPK model. Inputs to PBPK models comprise drug-independent and drug-dependent information. Organ mass or volume, blood flow, tissue composition, and the anatomical arrangement of the tissues and organs of the body are drug-independent components. Drug-dependent information includes partition coefficients, protein binding, PK properties, membrane permeability, enzymatic stability, and transporter-drug relationships. Since drug-independent components and many of the drug-dependent components are not required in the classical PK model, it is believed that the PBPK model contains richer informational content.

Besides data collection, the general procedure for developing PBPK models consists of three major steps [116]: 1) specification of whole body structure and the tissue structure, 2) writing differential equations, and 3) estimating parameters.

1.3.2.2. Specification of whole body structure and tissue structure

1.3.2.2.1. Specification of whole body structure. To establish a PBPK model for a specific compound, the whole body structure needs to be built to meet specific study purposes. It is important to decide which tissues/organs to include in the model. On one hand, the PBPK model should contain a large amount/number of tissues/organ that are important for drug ADME and tissues of special interest. On the other hand, for practical reasons, too many tissue/organ compartments not only increase the need for a lot of experimental data and literature information but also increase the difficulty of the mathematical calculations. Although currently there is no definite rule for selection of the tissues to be included, in general “core tissues” such as blood, liver, kidney, adipose, and tissues of interest are included in the model [116]. All the rest of the tissues can be lumped into “rapid equilibrating” or “slow equilibrating” compartments. When both a parental drug and its metabolite(s) are studied, separate PBPK models should be developed for both the drug and the metabolite(s). The models are then linked through the metabolism compartment (usually liver) with one part of the parent drug elimination output serving as input for the metabolite(s) [168, 169].

1.3.2.2.2. Specification of tissue structure. After a whole body structure is defined, the next step is to specify the structure of a specific tissue. In most cases, a simple perfusion limited model is used for the tissues (Figure 1-9A) [168]. The underlying assumption is that the drug distributes instantly in the tissues and there is no concentration gradient existing in the tissues. Although making this assumption oversimplifies the real situation, it has the major advantage of reducing the model complexity. When permeability limits the distribution of a drug within a tissue, a diffusion-limited model can be applied (Figure 1-9B) [168]. One tissue compartment can be divided further into two or three compartments. For a two compartment model in a
Figure 1-9. Structures and equations for individual organ models

A. Perfusion-limited model for non-eliminating organs. $V_T$ is the total volume of organ T; $Q_T$ is the blood flow to the organ; $C_T$ is the concentration of the drug in the organ; $R_T$ is the partition coefficient; $C_{Art}$ is the concentration of drug in arterial blood; $C_{T, Ven}$ is the venous effluent drug concentration ($C_{T, Ven} = C_T/R_T$). In this model, the measured tissue concentration is $C_T$.

B. Diffusion-limited model for non-eliminating organs. $C_{T,V}$ and $V_{T,V}$ represent the drug concentration and volume for the organ vascular space; $C_{T, EV}$ and $V_{T, EV}$ represent the corresponding terms for the extra-vascular space; $k_{V, EV}$ and $k_{EV, V}$ represent the drug transport rates between the vascular and extra-vascular spaces; $f_{ub}$ is the fraction of unbound drug in the vascular space. In this model, the measured tissue concentration is $C_{T, EV}$.

C. Perfusion-limited model for eliminating organ. $CL_{intT}$ represents the intrinsic clearance of the drug.

D. Diffusion-limited model for eliminating organs. $CL_{intT}$ represents the intrinsic clearance of the drug.

A | Perfusion limited model for non-eliminating organs

\[
V_T \frac{dC_T}{dt} = Q_T (C_{Art} - \frac{C_T}{R_T})
\]

B | Diffusion limited model for non-eliminating organs

\[
\begin{align*}
V_{T,Ev} \frac{dC_{T,Ev}}{dt} &= Q_T (C_{Art} - C_{Tv}) - k_{Ev,Ev} f_{Ev} C_{Tv} V_{T,Ev} + k_{Ev,Tv} C_{Tv} V_{T,Ev} \\
V_{T,Ev} \frac{dC_{T,Ev}}{dt} &= k_{Ev,Ev} f_{Ev} C_{Tv} V_{T,Ev} - k_{Ev,Tv} C_{Tv} V_{T,Ev}
\end{align*}
\]

C | Perfusion limited model for eliminating organs

\[
V_T \frac{dC_T}{dt} = Q_T (C_{Art} - \frac{C_T}{R_T}) - CLint_T C_T
\]

D | Diffusion limited model for eliminating organs

\[
\begin{align*}
V_{T,Ev} \frac{dC_{T,Ev}}{dt} &= Q_T (C_{Ev} - C_{Tv}) - k_{Ev,Ev} f_{Ev} C_{Tv} V_{T,Ev} + k_{Ev,Tv} C_{Tv} V_{T,Ev} \\
V_{T,Ev} \frac{dC_{T,Ev}}{dt} &= k_{Ev,Ev} f_{Ev} C_{Tv} V_{T,Ev} - k_{Ev,Tv} C_{Tv} V_{T,Ev} - CLint_T C_{T,Ev}
\end{align*}
\]
certain tissue, if the rate limiting step is assumed to happen in the capillary membrane, then the two compartments represent the vascular and extra-vascular compartments; if the rate limiting step is assumed to happen at the cell membrane, then the two compartments represent intra- and extra-cellular compartments. For the three compartment model in a certain tissue, it is assumed that both the capillary and cell membrane are rate-limiting steps for the drug. And the three compartments represent vascular, interstitial, and cellular compartments [116]. For eliminating organs, an eliminating function component is added to the model structures (Figure 1-9C, D) [168]. In reality, a whole body PBPK model can be composed of both perfusion- and diffusion-limited tissues. For example, the whole body PBPK model developed for 17-(allylamino)-17-demethoxygeldanamycin (17-AAG) contains both perfusion-limited organs (lung, brain, heart, spleen, kidney, and muscle) and diffusion-limited tissue (tumor) [168]. Whole body PBPK model developed for topotecan contains both perfusion-limited organs (lung, heart, muscle, adipose, skin, brain, liver, spleen, and gut) and diffusion-limited organs (kidney and testes) [170].

1.3.2.3. Writing differential equations. As in classic PK modeling, PBPK modeling equations are written following the mass balance equation. The difference is that the structure and components of the model represent real physiology. Examples of the equations are listed in Figure 1-9 [168]. During model development process, equations may be modified.

1.3.2.4. Estimating parameters. In order to estimate parameters, in addition to experimental data, physiological and anatomical data should be collected for modeling. However, obtaining reliable physiological parameters is not an easy task. Inaccurate physiological parameters used in the model will affect value of the final model. Among all the parameters, sensitivity analysis showed that blood flow is the most influential parameter [171]. Since various conditions such as anesthesia and stress can alter blood flow, using physiological data that are obtained under these conditions may bias the modeling results.

Because of the complexity of the PBPK model, PBPK parameter estimation usually is conducted from the open loop to closed loop (Figure 1-10). Blood concentration is first modeled using the empirical approach. The established blood PK model from the empirical approach is then used as a forcing function to each individual tissue. In this step, information such as blood flow, organ weight, and partition coefficient are used or estimated. Since individual tissues are still isolated, the model at this stage is open loop. Then the PBPK model is modified to a closed circulating system. Due to the computational challenges in modeling all the PBPK parameters at the same time, an open-loop approach followed by closed-loop approach is recommended [117].
Figure 1-10. Schematic structure of open loop and closed loop

In the open loop form, separate PBPK models were established for individual organs. Later in the closed loop form, all the modeled organs are linked together to form an anatomically integrated PBPK model. Q refers to blood flow to the lung (QL), organ 1 (Q1), organ 2 (Q2), and organ 3 (Q3). Input can be any site of the body.
1.3.3. Applications of PBPK model in drug discovery and development

1.3.3.1. Predicting human ADME and whole plasma PK profile using PBPK model. In early drug development stages, the measurements or simulation of drug-specific data—such as molecular weight, Log P (or Log D), pKa, plasma protein binding, and in vitro hepatic intrinsic clearance—can be incorporated into the PBPK model to predict animal and human ADME and facilitate candidate selection. Full plasma concentration-time profiles following different dosing schedules can be predicted by the PBPK models. For example, using the PBPK model, Jones et al. from F. Hoffmann-La Roche reported better prediction of human plasma concentration compared to commonly used allometric scaling (Dedrick approach) for compounds with distinct physicochemical and pharmacokinetic properties [119, 172]. In addition to the report from Jones et al., various approaches had been reported to predict ADME [119, 125, 173-180]. These prediction methods are generally complex, and the new trend and preferred approach is to perform prediction using commercially available PBPK software such as Simcyp, Gastroplus, and PK-Sim (mentioned in section 1.3.1.3).

Figure 1-11 represents the general scheme for predicting human PK data using this approach. Human concentration-time profiles under various dosing schedules can be generated when drug-specific data are provided. This can greatly simplify the ADME prediction process. In addition, animal PK data can be incorporated into the PBPK model to generate a human PK profile or to validate the IVIVE prediction. In this case, a PK parameter (such as clearance) estimated using the classic PK model is incorporated into the PBPK model to obtain a better prediction. More studies are needed to validate the predictive ability of the software. And more improvement is needed, especially for predicting oral absorption and for dealing with complex transporters involved in the PK process.

1.3.3.2. Drug distribution in target tissues and tumors. PBPK models have been used to understand and predict target tissue and tumor concentrations. In the oncology area, a PubMed search (09/12/2011) using “physiologically based pharmacokinetic model cancer” or “physiologically based pharmacokinetic model tumor” returned 153 papers. PBPK models had been used for antibody [155-158, 160-163, 181-183], liposomal [184], imaging agents [185-187], radioimmunodetection and radioimmunotherapy [188, 189], and small molecules such as topotecan [170], docetaxel [190, 191], gefitinib [192-194], moxifloxacin [195], capecitabine [196, 197], temozolomide [198, 199], genistein [200, 201], doxorubicin [136], 17-AAG [168], and methotrexate [202, 203]. General methods and principles of model development have been discussed in 1.3.1 and 1.3.2.

Partition coefficient is an important parameter in the PBPK model. Multiple methods for obtaining the partition coefficient values were applied as follows: 1. Estimation using PBPK model. 2. Estimation using non-compartmental modeling. Shah et al. reported a whole body PBPK model for topotecan [170]. In that model, a tissue to
Peff, jejunum permeability; AUC, area under the concentration vs. time curve; B/P, blood to plasma ratio; Cmax, maximum concentration; CL, clearance; CLint, intrinsic clearance; CLr, renal clearance; DDI, drug–drug interaction; EC50 or IC50, concentration causing half of the maximal effect of induction or inhibition; Emax or Imax, maximum effect of induction or inhibition; F, bioavailability; Fa, fraction absorbed; Fg, bioavailability in the gut; Fh, bioavailability in the liver; fu,p, unbound fraction in plasma; γ, Hill coefficient; Jmax, maximum rate of transporter-mediated efflux/uptake; Ka, first-order absorption rate constant; Kd, dissociation constant of drug–protein complex; Ki, reversible inhibition constant; KI, apparent inactivation constant, concentration causing half of the maximal inactivation; kinact, apparent maximum inactivation rate constant; Km, Michaelis–Menten constant, substrate concentration causing half of the maximal reaction or transport; Kp, tissue-to-plasma partition coefficient; LogP, logarithm of the octanol–water partition coefficient; MOA, mechanism of action; PD, pharmacodynamics; PK, pharmacokinetics; PopPK, population pharmacokinetics; V, volume of distribution; Vmax, maximum rate of metabolite formation.

plasma partition coefficient for each tissue was first estimated using the ratio of AUC (estimated from non-compartmental modeling) for the tissue to the ratio in plasma. These values were then fixed and applied to the whole body PBPK model. 3. *In vitro* measurements. Bradshaw-Pierce et al. reported a PBPK model for docetaxel [204]. The PBPK model for docetaxel was developed incorporating specific binding of docetaxel to intracellular components, liver metabolism, biliary elimination, and fecal and urinary excretion. Tissue/blood partition coefficients were determined *in vitro* [204, 205]. Docetaxel was incubated with minced tissue at 37°C with gentle shaking for 24 h. Samples were centrifuged to separate tissue from saline. Docetaxel concentration was measured in both the saline and tissue layer. The partition coefficient was determined by the ratio of docetaxel concentrations in the tissue layer to the saline layer. These values were fixed and applied to the whole body PBPK model. 4. Calculation. Sung et al. reported a PBPKPD model for UTF (5-FU, tegafur, and uracil) [206]. Partition coefficients for tissues were calculated based on the n-octanol-water partition coefficient developed by Poulin and Theil [207, 208]. The estimation of partition coefficient for tumor was adjusted during model development. Once a PBPK model is established, scaling the concentration from one species to another can be done by changing the physiological information but keeping the model structure and assuming identical partition coefficients between species [168, 192, 198].

**1.3.3.3. Predicting the risk for drug-drug interactions.** Drug-drug interactions can cause serious toxicities or loss of efficacy. Due to the fact that many drugs are substrates and/or inhibitors of various drug metabolizing enzymes and efflux transporters, the degree of drug-drug interaction is a concern when drugs are used in combination. To understand the potential for a drug-drug interaction for a certain drug, evaluations are usually conducted in clinical trials or in preclinical settings. Now, by combining the physiological information and the drug metabolism enzyme/transporter inhibition/or induction knowledge obtained *in vitro* experimentally, the PBPK model can be used to simulate and predict drug-drug interaction in humans. Multiple PBPK model applications for predicting drug-drug interactions have been reported [130, 143-150]. For example, Johnson et al. built a pediatric PBPK (P-PBPK) model to predict drug concentration-time profiles and drug-drug interactions in children [130]. Using Simcyp Pediatric ADME Simulator, concentration time profile was simulated for a 2-year-old child prescribed with CYP3A4 inducer, CYP3A4 inhibitor, and substrates with complex dosing schedules.

The FDA reported example cases of using a PBPK model during investigational new drugs (INDs) and new drug applications (NDAs) reviews between July 2008 and June 2010 [149]. In these cases, the PBPK approach had been used to address multiple specific regulatory questions, such as whether a drug-drug interaction study can be ruled out even though the ratio of *in vivo* inhibitor concentration I (maximum plasma concentration at the highest proposed dose) to the *in vitro* inhibition constant (Ki) (I/Ki) is > 0.1. The software used in the study included GetData 2.24 (for digitizing concentration-vs.-time profiles), WinNonlin 5.2, NONMEM (for PK parameter estimation), and PBPK simulators (Simcyp, PK-Sim, Gastroplus, and SAAMII). At the same time, it should be mentioned that the PBPK model used in this case relies largely on
in vitro and in silico data. Some important parameters needed may not be available in earlier drug development stage. As the compound progresses to later stage, the PBPK model can be refined incorporating the available data.

### 1.3.4. Limitation of PBPK modeling

So far, in comparison to the classical PK model, the application of the PBPK approach is still limited despite significant potential. Developing a PBPK model is demanding because of the investment of large amount of time and effort to obtain information needed to establish the model. In addition, data required to establish the model are not always available. Variations in physiological parameters obtained from the literature, our current limited knowledge about the underlying mechanism about drug ADME process, and wrong assumptions can all affect the quality of the model. The PBPK model is usually structurally complex and much more methodologically and computationally challenging compared to the classic PK model. Furthermore, PBPK model is not as mature as conventional PK model.

Despite all of the limitations of the PBPK model mentioned above, the PBPK area is developing rapidly. The knowledge and methods for building the model are continuously accumulating and improving. The PBPK model breaks the limit of conventional PK by building a model and understanding the PK process based on the real physiological and anatomical system, and by incorporating rich knowledge learned over the years about effect of drug dependent and independent factors on PK. In the drug discovery and development area, pharmaceutical companies are acquiring PBPK modeling software for predicting human PK based on in vitro and/or animal in vivo PK data. Also, the regulatory authority has started appreciating the benefit of the PBPK model and applying the PBPK model to address drug regulatory review questions. At last, but not least, the PBPK model will be applied with increased frequency in the toxicology and oncology areas where understanding target tissue or tumor concentration is important yet data are usually inaccessible in the human.

### 1.4. Summary

Reactivating the p53 pathway is regarded as an appealing nongenotoxic approach for treating tumors with wild-type p53. Unlike adult cancers, pediatric malignancies usually retain a high percentage of wild-type p53 status at diagnosis. Using neuroblastoma as an example, 98% of neuroblastoma tumors retain wild-type p53 at diagnosis [209, 210]. Even in relapse neuroblastoma, a majority of the tumors are still p53 wild-type [211]. Nutlin-3a is a small molecule inhibitor that targets the MDM2/MDMX-p53 interaction. It is currently under pre-clinical investigation in a variety of solid tumor and leukemia models, and has shown promising in vitro and in vivo activities. Like most other anticancer drugs, it is most likely that nutlin-3 will be used in combination. Studies have suggested synergistic effects when nutlin-3 and other chemotherapeutic drugs were co-administered, independent of p53 status, by enhancing
the ability of anticancer drugs to activate apoptosis or by reversing P-gp-mediated drug resistance. ABC transporters play important roles in drug resistance and ADME processes. The effect of nutlin-3a on BCRP had not been reported before.

Understanding the pharmacokinetics of a compound is critical in preclinical drug development. An understanding of the systemic disposition of nutlin-3a, as well as the distribution to target tissue or tumor sites, will provide a rational basis for the selection of dosage regimens for preclinical models. To date, the pharmacokinetics of nutlin-3a has not been reported. Whole-body PBPK model, which is based on anatomical compartments and blood flow, is an excellent tool to describe and predict drug concentration not only in blood but also in target tissues.

1.5. Specific Aims

The objective of specific aim 1 was to assess the effect of nutlin-3a on cell viability, both as a single agent and in combination with one of the commonly used anti-neuroblastoma agents, topotecan.

The objective of specific aim 2 was to investigate whether nutlin-3a inhibits BCRP, thus sensitizing cells to enhanced killing by anti-cancer drugs that are BCRP substrates. Based on initial observations, we performed a series of studies to investigate the effect of nutlin-3a treatment on BCRP expression and function comprehensively [212].

The objective of specific aim 3 was to perform pharmacokinetic studies and develop a PBPK model describing the disposition of nutlin-3a in plasma and tissues, including adipose, adrenal gland, bone marrow, brain, liver, lung, intestine, muscle, retina, spleen, and vitreous fluid. The PBPK model was used to perform simulations, which—in combination with in vitro cell sensitivity data—provided rationale for choosing dosing regimens for mouse models of common childhood cancers, including retinoblastoma, neuroblastoma, rhabdomyosarcoma, and acute lymphoblastic leukemia (ALL) [213].
CHAPTER 2. MDM2 ANTAGONIST NUTLIN-3A SYNERGISTICALLY INHIBITS NEUROBLASTOMA CELL GROWTH WITH TOPOTECAN

2.1. Introduction

Neuroblastoma [214] is an embryonic malignancy of the sympathetic nervous system. It is the second most common extra-cranial solid tumor of childhood [215] and the most common cancer of infancy [216]. While other childhood cancers have experienced an improved cure rate over the past decades, high risk neuroblastoma is still one of the most difficult tumors to treat, with less than 30% long-term survival despite aggressive multi-modal therapy including surgery, radiation, high-dose chemotherapy, and stem cell transplantation [217].

To provide a new therapy for neuroblastoma, effective molecular targets should be selected. One of the targets that we studied is p53. p53 plays a central role in regulating cell cycle arrest, senescence, and apoptosis. The p53 gene is mutated in half of all tumors, and in those tumors that retain wild-type p53, p53 function may also be suppressed by its negative regulators. As a key negative regulator of p53 [218], oncogene MDM2 effectively impairs p53 stability and activity. Strong evidence for the relationship between MDM2 and p53 has made inhibition of the MDM2-p53 interaction an attractive target for reactivating p53.

One crucial point of neuroblastoma is that 98% of neuroblastoma tumors retain wild-type p53 at diagnosis [209, 210]. Even in relapse neuroblastoma, a majority of the tumors are still p53 wild-type [211]. Although many mechanisms for inactivation of p53 pathway had been proposed [211, 219], several studies indicate that the p53 downstream pathway is intact in neuroblastoma and can be activated in response to DNA damage [220]. These data suggest that regardless of many other unclear possible mechanisms of p53 inactivation, p53 downstream pathway is intact, which encouraged us to restore the function of p53 by directly stabilizing p53 protein using the MDM2-p53 interaction inhibitor to treat neuroblastoma.

The results of preclinical studies have shown that the MDM2 antagonist nutlin-3a is a promising agent to reactivate p53. Nutlin-3a has demonstrated anti-tumor efficacy in preclinical models of human osteosarcoma, retinoblastoma, and KSHV lymphoma with wild-type p53 [17, 24, 25, 39-41]. In neuroblastoma, studies have found that MDM2 antagonist nutlin-3a is effective both in vitro [36, 221, 222] and in vivo [43].

Several studies suggested that when nutlin-3a was combined with anti-cancer agents, synergistic growth inhibition was observed [31, 35, 37]. In neuroblastoma, nutlin-3 synergized (or sensitized) neuroblastoma cells to chemotherapeutic agents (R)-roscovitine [36], cisplatin [221], etoposide [221], and doxorubicin [223, 224]. So far, combining nutlin-3 with topotecan, a commonly used anti-neuroblastoma agent, has not been reported in neuroblastoma research.
In this study, we reported that nutlin-3a synergistically inhibited neuroblastoma cell growth with topotecan. To explore a possible mechanism, we studied the effect of nutlin-3a on the expression of efflux transporter P-gp.

2.2. Materials and Methods

2.2.1. Reagents

Nutlin-3a were synthesized in the Department of Chemical Biology at St. Jude Children’s Research Hospital, Memphis, TN, and were solubilized in dimethyl sulfoxide (DMSO) (ATCC, Manassas, VA) to a final concentration of 20 mM. Topotecan hydrochloride was purchased from GlaxoSmithKline (Philadelphia, PA) and solubilized to a final concentration of 20 mM in sterile water. CellTiter 96® AQueous MTS assay reagents were purchased from Promega (Madison, WI).

2.2.2. Cell culture

Neuroblastoma cell lines NB1691, NB1643, NBEBC1, SHSY-5Y, SKNAS, and SKNBE2C were cultured in antibiotic-free RPMI-1640 medium (Cellgro®, Manassas, VA) containing 10% fetal bovine serum (FBS) (Invitrogen, Carlsbad, CA) and 1% L-Glutamine (Invitrogen, Carlsbad, CA) at 37°C in 5% CO₂.

2.2.3. Cell viability assay

Briefly, cells were seeded in 100 µl phenol red-free RPMI-1640 medium (Cellgro®, Manassas, VA) in 96-well plates and allowed to attach overnight. Cells were treated with increasing concentrations of nutlin-3a alone or increasing concentrations of topotecan in combination with nutlin-3a for 6 hours. After 6 hours, topotecan was washed off and cells were incubated with or without nutlin-3a for additional 18 hours. Cell viabilities were tested by MTT assay or CellTiter 96® AQueous MTS assay following the manufacturer's protocol. IC₅₀ values were calculated using ADAPT 5 (Biomedical Simulations Resources, Los Angeles, CA) [225].

2.2.4. Synergy study

Synergism, additive effects, and antagonism were assessed using the median effect method developed by Chou [226]. The combination index (CI) values at non-fixed nutlin-3a/topotecan concentration ratios were calculated using the commercially available software CalcuSyn 2.1 (Biosoft, Cambridge, United Kingdom). CI values < 1.0 indicate synergism, CI values = 1.0 indicate additive effect, and CI values > 1.0 indicate antagonism [226].
2.2.5. **qRT-PCR**

Cells were treated with 10 μM nutlin-3a for 24 hours. Total RNA samples were extracted using Qiagen RNeasy® Plus Mini Kit (Valencia, CA) following manufacturer's instructions. RNA concentrations were quantified using Beckman DU-600 spectrophotometer (Beckman Instruments Inc., Fullerton, CA). Genomic DNA elimination, RT reaction, and cDNA synthesis reaction were performed using the RT² First Strand Kit (SABiosciences, Frederick, MD) following the manufacturer's protocol.

Primers used for RT-PCR analysis were synthesized by the Hartwell Center at St. Jude. Children’s Research Hospital. For *MDRI*, the forward primer sequence was CCATCATTGCAATAGCAGG, and the reverse primer sequence was TGTTCAAACTTCTGCTCCTGA. For GAPDH control, the forward primer sequence was AAGGACTCATGACCACAGTCCAT, and the reverse primer sequence was CCATCAGCCACAGTTTCC. SYBR Green qPCR Master Mix was purchased from SABiosciences (Frederick, MD). QRT-PCR reactions were performed on ABI Prism® 7700 Sequence Detection System (Applied Biosystems, Foster City, CA). The reactions were conducted under the following conditions:

<table>
<thead>
<tr>
<th>Time</th>
<th>Temp</th>
<th>Cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td>15 min</td>
<td>95 °C</td>
<td>1 cycle</td>
</tr>
<tr>
<td>18 sec</td>
<td>95 °C</td>
<td>40 cycles</td>
</tr>
<tr>
<td>30 sec</td>
<td>52 °C</td>
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</tr>
<tr>
<td>15 sec</td>
<td>95 °C</td>
<td>1 cycle</td>
</tr>
<tr>
<td>15 sec</td>
<td>60 °C</td>
<td></td>
</tr>
<tr>
<td>15 sec</td>
<td>95 °C</td>
<td></td>
</tr>
</tbody>
</table>

2.2.6. **Western blots**

Cells were lysed in M-PER Mammalian Protein Extraction Reagent (Thermo Scientific, Rockford, IL) following manufacturer’s instructions. Protein concentrations were determined using the BCA protein assay (Thermo Scientific Rockford, IL). 8-16 μg of proteins were resolved by SDS PAGE gels (4–12% gradient Bis/Tris NuPage gels) (Invitrogen, Carlsbad, CA) with MOPS SDS running buffer (Invitrogen, Carlsbad, CA) before transferring onto 0.45 μm pore size InvitrolonTM PVDF membranes (Invitrogen, Carlsbad, CA). P-gp was detected using the mouse monoclonal antibody clone C-219 (Alexis Biochemicals, San Diego, CA). β -actin (AC-15, Sigma–Aldrich, St. Louis, MO) was used as the loading control.
2.2.7. Statistical analysis

All data were expressed as mean ± standard deviation. Data were analyzed for statistical significance using Student’s t-test. Differences with p < 0.05 were considered statistically significant.

2.3. Results

2.3.1. Effect of single agent nutlin-3a on the cell viability of neuroblastoma cell lines

p53 wild-type NB1691, NB1643 and p53 mutant SKNBE2C cells were incubated with increasing concentrations of nutlin-3a alone for 24 hours. IC\textsubscript{50} values were determined by MTT assay (Figure 2-1). The IC\textsubscript{50} values of nutlin-3a in p53 wild-type cells NB1691 and NB1643 were 3.99 (± 0.12) μM and 4.29 (± 0.20) μM, respectively. p53 mutant SKNBE2C cells were resistant to nutlin-3a treatment, with IC\textsubscript{50} value of 26 (± 295.62) μM.

2.3.2. Combination of nutlin-3a with topotecan in NB1691 cells

NB1691 cells were incubated with increasing concentrations of topotecan in combination with DMSO control or 2μM nutlin-3a for 6 hours. After 6 hours, cells were washed with drug-free media. Media containing either DMSO control or 2μM nutlin-3a was added back to the wells for additional incubation of 18 hours. IC\textsubscript{50} values were determined by MTT assay (Table 2-1). Without combination, the IC\textsubscript{50} value of topotecan in NB1691 cells was 8.39 (± 1.04) μM. Combination of topotecan with nutlin-3a resulted in a 4.7-fold reduction of IC\textsubscript{50} to 1.79 (± 0.29) μM.

2.3.3. Synergistic effect of nutlin-3a in combination with topotecan

To study the interaction of nutlin-3a and topotecan in NB1691 cells, NB1691 cells were incubated with various concentrations of nutlin-3a and topotecan. Cells were treated with nutlin-3a and topotecan for 6 hours. After 6 hours, cells were washed with drug-free media. Media containing various concentrations of nutlin-3a was added back to the wells for additional incubation of 18 hours. IC\textsubscript{50} values were determined by MTS assay.

The CI values at non-fixed nutlin-3a/topotecan concentration ratios were calculated using CalcuSyn 2.1. The calculated CI values were listed in Table 2-2. Fractional effect is defined as the fraction of cells affected by nutlin-3a and topotecan combination. A fractional effect value of 0 indicates no inhibition, and a fractional effect value of 1 indicates 100% inhibition of cell viability. When topotecan concentrations were less than 25 μM, CI values were generally < 1, indicating a synergistic effect. When
Figure 2-1. Effect of single agent nutlin-3a on the cell viability of neuroblastoma cell lines

p53 wild-type NB1691, NB1643 and p53 mutant SKNBE2C cells were incubated with increasing concentrations of nutlin-3a alone for 24 hours. IC\textsubscript{50} values were determined by MTT assay in 5 replicates. IC\textsubscript{50} values were calculated using ADAPT 5. Values are presented as mean ± SD. Data are representative of one experiment.

Table 2-1. Combination of nutlin-3a with topotecan in NB1691 cells

<table>
<thead>
<tr>
<th>Treatment</th>
<th>IC\textsubscript{50} (µM)</th>
<th>SD</th>
<th>Confidence interval (95%) (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Topotecan</td>
<td>8.39</td>
<td>1.04</td>
<td>[6.32, 10.46]</td>
</tr>
<tr>
<td>Nutlin-3a +topotecan</td>
<td>1.79</td>
<td>0.29</td>
<td>[1.21, 2.38]</td>
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</table>

NB1691 cells were incubated with increasing concentrations of topotecan in combination with DMSO control or 2 µM nutlin-3a for 6 hours. After 6 hours, cells were washed with media. Media containing either DMSO control or 2µM nutlin-3a was added back to the wells for additional incubation of 18 hours. IC\textsubscript{50} values were determined by MTT assay. Combination of topotecan with nutlin-3a resulted in a 4.7-fold reduction of IC\textsubscript{50} to from 8.39 (± 1.04) µM to 1.79 (± 0.29) µM. IC\textsubscript{50} values were calculated using ADAPT 5. Data are representative of one experiment.
Table 2-2. Synergistic effect of nutlin-3a in combination with topotecan

<table>
<thead>
<tr>
<th>Topotecan (μM)</th>
<th>Nutlin-3a (μM)</th>
<th>Fa</th>
<th>Combination index (CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.39</td>
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<td>0.46</td>
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</tr>
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<td>2.50</td>
<td>0.46</td>
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</tr>
<tr>
<td>0.39</td>
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<td>0.46</td>
<td>1.15</td>
</tr>
<tr>
<td>0.39</td>
<td>10.00</td>
<td>0.68</td>
<td>0.71</td>
</tr>
<tr>
<td>0.39</td>
<td>20.00</td>
<td>0.67</td>
<td>1.56</td>
</tr>
<tr>
<td>0.78</td>
<td>1.25</td>
<td>0.47</td>
<td>0.46</td>
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<td>0.49</td>
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<td>0.79</td>
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<tr>
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<td>6.56</td>
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<td>20.00</td>
<td>0.75</td>
<td>1.46</td>
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Table 2-2. (Continued)

<table>
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<tr>
<th>Topotecan (µM)</th>
<th>Nutlin-3a (µM)</th>
<th>Fa</th>
<th>Combination index (CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>100.00</td>
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<td>0.63</td>
<td>4.91</td>
</tr>
<tr>
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<td>0.63</td>
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</tr>
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<td>5.26</td>
</tr>
<tr>
<td>100.00</td>
<td>10.00</td>
<td>0.62</td>
<td>6.34</td>
</tr>
<tr>
<td>100.00</td>
<td>20.00</td>
<td>0.64</td>
<td>5.87</td>
</tr>
</tbody>
</table>

NB1691 cells were incubated with various concentrations of nutlin-3a and topotecan. IC₅₀ values were determined by MTS assay. The CI values at non-fixed nutlin-3a/topotecan concentration ratios were calculated using commercially available software Calcusyn 2.1. CI values < 1.0 indicate synergism, CI values = 1.0 indicate additive effect and CI values > 1.0 indicate antagonism. Data are representative of one experiment.
topotecan concentrations were higher than 25 µM, CI values were > 1, indicating antagonism.

2.3.4. MDRI gene expression change after nutlin-3a treatment

Since wild-type p53 was reported to repress the MDRI promoter [97-99], the synergism observed in NB1691 may be explained by the reduction of MDRI gene expression. To evaluate the effect of nutlin-3a on the MDRI gene expression, we performed qRT-PCR in a panel of neuroblastoma cell lines after 10 µM nutlin-3a treatment for 24 hours. Compared to control cells, a 3-fold reduction of MDRI gene expression was observed in MDM2-amplified NB1691 cells. In contrast, a 1.54-fold increase of MDRI gene expression was observed in p53 mutant SKNBE2C cells. There were no changes in the gene expression of MDM2 non-amplified p53 wild-type cells SHSY5Y, NB1643, and NBEBC1 cells and p53 null SKNAS cells. Results in Figure 2-2 represent pooled data of three independent experiments.

2.3.5. P-gp protein expression after nutlin-3a treatment

In addition to evaluating MDRI expression at the message level, P-gp expression at the protein level was evaluated using western blot. Results in Figure 2-3 suggest reduced protein expression of P-gp in NB1691 and NBEBC1 cells. No increased P-gp protein expression was observed in SKNBE2C cells.

2.4. Discussion

Our preliminary studies showed that targeting MDM2-p53 interaction using nutlin-3a reduced cell growth in neuroblastoma cells. p53 wild-type cells were much more sensitive to nutlin-3a treatment compared to p53 mutant cells. When nutlin-3a was combined with topotecan, a synergistic effect was observed when topotecan concentrations were < 25 µM. When topotecan concentrations were > 25 µM, the nutlin-3a and topotecan combination caused antagonistic effect. This is the first study looking at nutlin-3a and topotecan interaction in neuroblastoma cell lines. Synergism has been reported in other tumor types, such as retinoblastoma [24] and intraocular melanoma [227], when nutlin-3a and topotecan were combined. Topotecan concentrations tested in those two studies did not include levels > 25 µM.

To investigate possible mechanism of synergy, we performed qRT-PCR and western blot and found reduced MDRI expression at both the message level and protein level in MDM2-amplified NB1691 cells. In MDM2 non-amplified NBEBC1 cells, although no MDRI gene expression change was observed, western blot data showed reduced expression of P-gp protein. Since topotecan is a substrate of P-gp, synergism observed may be partly explained by reduced P-gp expression. However, it should be
Figure 2-2.  *MDRI* gene expression change after nutlin-3a treatment

*MDRI* gene expression changes after 10 μM nutlin-3a treatments were evaluated using qRT-PCR in a panel of neuroblastoma cell lines. Values are presented as mean ± SD. Data are representative of three independent experiments with triplicates in each experiment.

Figure 2-3.  P-gp protein expression change after nutlin-3a treatment

Western blot from cell lysates collected after 24-hr nutlin-3a treatment (20 μM) probed using antibody c219 against P-gp. β-actin was used as loading control. Data are representative of one experiment.
mentioned that this study was preliminary, and more independent experiments are needed to support the change of P-gp expression at the protein level.

As we were planning to conduct further research to study the effect of nutlin-3a on P-gp function, Michaelis et al. in the meantime reported that nutlin-3a inhibited P-gp function and reversed vincristine resistance in neuroblastoma and rhabdomyosarcoma cells independent of p53 status [228]. The mechanism of inhibition proposed in that study was competitive inhibition. In the current study, we evaluated a different mechanism of action of nutlin-3a on P-gp. Instead of competitive inhibition, nutlin-3a inhibited the expression of P-gp.
CHAPTER 3. MDM2 ANTAGONIST NUTLIN-3A REVERSES MITOXANTRONE RESISTANCE BY INHIBITING BREAST CANCER RESISTANCE PROTEIN MEDIATED DRUG TRANSPORT*

3.1. Introduction

Pre-clinical investigations of the utility of nutlin-3 treatment of cancer cells have focused primarily on the consequences of p53 reactivation in cells due to disruption of the MDM2/p53 interaction. Nutlin-3 is a racemic mixture of nutlin-3a (active enantiomer) and nutlin-3b (inactive enantiomer) with nutlin-3a having 150-fold more affinity to MDM2 [17]. Indeed, single agent nutlin-3 treatment has shown anti-cancer efficacy in xenograft models of solid tumors, including osteosarcoma, prostate cancer, KSHV lymphomas, retinoblastoma, and neuroblastoma [17, 24, 25, 40, 43]. Recently, other effects of nutlin-3 treatment have been reported, including anti-angiogenic effects [214, 229, 230] and radiosensitization of cancer cells under low oxygen conditions [29]. Furthermore, nutlin-3 has been reported to sensitize cancer cells to co-treatment with selected anti-cancer drugs, independent of p53 status, by enhancing the ability of anticancer drugs to activate apoptosis [223], and also by reversing P-gp mediated MDR [228]. Understanding the mechanism behind this nutlin-3 sensitization of resistant cancer cells would significantly enhance the use of nutlin-3 in combination with other anti-cancer drugs in a broad range of tumor types.

Drug-resistance is a major obstacle in the treatment of cancer, and ATP-binding cassette (ABC) transporters play an integral role in the development of multi-drug resistance [231]. ABC transporters utilize the energy of ATP hydrolysis to pump anti-cancer agents out of the cell, thus reducing the intracellular drug concentration. Recently, Michaelis et al. observed that nutlin-3 can interfere with the function of the ABC transporters P-pg and MRP1 [228]. Nutlin-3 treatment reversed drug resistance in neuroblastoma and rhabdomyosarcoma cells over-expressing these transporters in vitro when combined with cytotoxic drugs that are P-gp and MRP1 substrates. These data suggest that nutlin-3 functionally inhibits the action of drug efflux proteins, thereby sensitizing cells to treatment with cytotoxic agents that are substrates of these efflux proteins.

BCRP belongs to the ABC transporter family. Although it is possible that nutlin-3a may modulate the activity of BCRP, so far, the effect of nutlin-3a on BCRP has not been reported. The present study investigates whether nutlin-3a inhibits BCRP, thus sensitizing cells to enhanced killing by anti-cancer drugs that are BCRP substrates. Using MTS assays, we determined that nutlin-3a reverses resistance to the BCRP substrate mitoxantrone. Combination index calculations indicated synergism when nutlin-3a was

used in combination with the anticancer agent mitoxantrone, a BCRP substrate, in osteosarcoma cells over-expressing BCRP. Based on these observations, we performed a series of studies to comprehensively investigate the effect of nutlin-3a treatment on BCRP expression and function. Our studies strongly suggest that nutlin-3a inhibits BCRP efflux and can reverse BCRP-related drug resistance, but is not a BCRP substrate.

3.2. Materials and Methods

3.2.1. Reagents

Nutlin-3a and nutlin-3b were synthesized in the Department of Chemical Biology at St. Jude Children’s Research Hospital, Memphis, TN and were solubilized in dimethyl sulfoxide (DMSO) (ATCC, Manassas, VA) to a final concentration of 30 mM. The chemical structure of nutlin-3 has been published previously [17]. Hoechst 33342 and G-418 (Geneticin®) were purchased from Invitrogen (Carlsbad, CA). Mitoxantrone and Ko143 were purchased from Sigma-Aldrich (St. Louis, MO). Fumitremorgin C (FTC) was purchased from Alexis Biochemicals (San Diego, CA).

3.2.2. Cell culture

Saos-2 (human osteosarcoma) cells stably transfected with human wild-type (Arg482) BCRP or control vector pcDNA3.1 were generously donated by Dr. John Schuetz (St. Jude Children’s Research Hospital, Memphis, TN) [62, 232]. MDCK II-pCDNA3.1 and MDCK II-BCRP cells were generously donated by Dr. Mark Leggas (University of Kentucky, Lexington, KY). Cells were cultured in complete Dulbecco’s Modified Eagle’s Medium (DMEM) (Invitrogen, Carlsbad, CA) containing 10% fetal bovine serum (FBS) (Invitrogen, Carlsbad, CA), 1% L-glutamine (Invitrogen, Carlsbad, CA), and maintained with G418 (0.5 mg / ml) (Invitrogen, Carlsbad, CA). Cells were cultured in G418-free complete media at the time of seeding for individual experiments. Hank’s balance salt solution (HBSS) and HEPES buffer were purchased from Invitrogen (Carlsbad, CA).

3.2.3. Cell viability assay (MTS)

Cells were seeded in 100 μl phenol red-free medium (Invitrogen, Carlsbad, CA) in 96-well plates and allowed to attach overnight. Cells were treated with increasing concentrations of nutlin-3a alone or increasing concentrations of mitoxantrone in combination with 0 μM, 20 μM or 50 μM nutlin-3a for 24 hours. Cell viabilities were tested by the CellTiter 96® AQueous MTS assay (Promega, Madison, WI) following the manufacturer's instructions. IC50 values were calculated using ADAPT 5 (Biomedical Simulations Resources, Los Angeles, CA) [225].
3.2.4. Median effect analysis

To characterize the interaction between mitoxantrone and nutlin-3a in Saos-2-BCRP and Saos-2-pcDNA3.1 cells, data were analyzed using the median effect method developed by Chou [226]. The combination index (CI) values at non-fixed nutlin-3a/mitoxantrone concentration ratios were calculated using the commercially available software Calcusyn 2.1 (Biosoft, Cambridge, United Kingdom). CI values < 1.0 indicate synergism, CI values = 1.0 indicate additive effect and CI values > 1.0 indicate antagonism [226].

3.2.5. Intracellular accumulation and efflux of mitoxantrone by confocal imaging

Vector control and BCRP expressing Saos-2 cells were seeded on 35 mm glass bottom dishes (MatTek Corporation, Ashland, MA) and allowed to attach for 36 hours. Cells were pre-incubated with nutlin-3a for 15 minutes in DMEM+ (DMEM with 2% FBS, 1 mM HEPES buffer) at 37°C. Mitoxantrone (1 μM) was added and cultures were incubated for an additional 1 hour. Cells were washed with ice cold HBSS+ (HBSS with 2% FBS, 1 mM HEPES buffer) containing nutlin-3a and intracellular accumulation was measured using confocal imaging. An Eclipse C1si confocal, configured on an Eclipse TE2000 microscope (Nikon, Melville, NY) with a Plan Fluor 40 × NA 1.3 lens was used. Excitation was from a 642 nm diode laser, and the emission was collected through a 675/50 nm bandpass filter.

3.2.6. Hoechst 33342 dye accumulation and efflux studies by flow cytometry

Hoechst 33342 dye was used as a BCRP substrate. Vector control and BCRP over-expressing Saos-2 and MDCKII cells were cultured to 60-70% confluence. Single cell suspensions were pre-incubated in DMEM + with nutlin-3a, nutlin-3b or FTC at varying concentrations for 15 minutes at 37°C. Hoechst 33342 dye was then added to a final concentration of 5 μg/ml and cells were incubated at 37°C for 1 hour. Cells were pelleted and resuspended in ice cold HBSS+ containing nutlin-3a, nutlin-3b or FTC. Intracellular Hoechst 33342 fluorescence signals were detected by a 440/40 nm band pass filter with UV laser excitation and the data were collected and analyzed using a BD LSRII flow cytometer (BD, San Jose, CA). Data were processed as previously described [62]. Propidium iodide (PI) (Roche Applied Science, Mannheim, Germany) was added as a marker to label the non-viable cells. Only events from viable cells were used for data analysis.

3.2.7. Intracellular accumulation and efflux of Hoechst 33342 by widefield imaging

Vector control and BCRP expressing Saos-2 cells were seeded on 35 mm glass bottom dishes and allowed to attach for 36 hours. Cells were pre-incubated with nutlin-3a
or nutlin-3b in DMEM+ for 15 minutes at 37°C. Hoechst 33342 (1 μg / ml) was added and cultures were incubated for an additional 1 hour. Cells were washed with ice cold HBSS+ containing nutlin-3a or nutlin-3b and intracellular accumulation was measured using widefield fluorescence imaging. Hoechst 33342 imaging was performed on a Nikon Eclipse TE2000 microscope with a Plan Fluor 40 × NA 0.6 lens and a standard DAPI filter set.

3.2.8. Western blots

Total cellular protein was extracted from cell pellets and protein concentrations were determined by BCA assay (Thermo Scientific, Rockford, IL). Proteins were resolved by SDS PAGE [4-12% gradient Bis/Tris NuPage gels (Invitrogen, Carlsbad, CA)] with MOPS SDS running buffer (Invitrogen, Carlsbad, CA) before transferring onto InvitrolonTM PVDF membranes (Invitrogen, Carlsbad, CA). BCRP was detected using the rat monoclonal BXP-53 (Alexis Biochemicals, San Diego, CA). P-gp was detected using the mouse monoclonal antibody clone C-219 (Alexis Biochemicals, San Diego, CA). Beta-actin (AC-15, Sigma-Aldrich, St. Louis, MO) was used as the loading control.

3.2.9. Flow cytometry for BCRP

PE conjugated mouse anti-BCRP (MAB4155P) (Millipore, Billerica, MA) antibody was used to detect protein expression by flow cytometry. Cells were treated with 50 μM nutlin-3a for 1 hour, harvested, and then 1.0 × 10⁶ cells were resuspended in 100 μL BD Fc BlockTM (BD Pharmingen, San Diego, CA) and incubated on ice for 30 minutes. Cells were washed once and then stained with the primary antibody (anti-BCRP 10 μg / ml) at room temperature for 30 minutes. Cells were again washed and resuspended in a final volume of 0.5 ml. Cells were counterstained with 4’-6-Diamidino-2-phenylindole (DAPI) (Invitrogen, Carlsbad, CA) and analyzed using a BD LSRII flow cytometer (BD, San Jose, CA).

3.2.10. BCRP localization

Localization of BCRP was evaluated using confocal imaging analysis. Cells were grown on 35 mm glass bottom dishes and treated with 50 μM nutlin-3a for 90 minutes. After incubation, cells were washed with HBSS and fixed with 4% paraformaldehyde (Polysciences, Warrington, PA) for 15 minutes at room temperature. After 3 HBSS washes, cells were permeabilized using 0.2 % Triton X-100 (Sigma-Aldrich, St. Louis, MO) diluted in HBSS for 5 minutes then washed in HBSS. After 3 HBSS washes, Image-iT™ FX signal enhancer solution (Invitrogen, Carlsbad, CA) was applied and the dishes were incubated at room temperature for 30 minutes. After 3 additional HBSS washes, mouse monoclonal antibody against BCRP (BXP-21, Alexis Biochemicals, San Diego, CA) (1:1000 diluted in HBSS + 2% bovine serum albumin) was added and cells were incubated for an additional 60 minutes at room temperature. Cells were then stained
with Alexa Fluor® 555 goat anti-mouse IgG secondary detection conjugate (1:500) (Invitrogen, Carlsbad, CA) at room temperature for 90 minutes. Finally, ProLong® Gold-DAPI anti-fade reagent (Invitrogen, Carlsbad, CA) was added and wells were cover-slipped. BCRP localization was assessed using confocal imaging. A Nikon Eclipse C1si confocal configured on an Eclipse TE2000 microscope with a Plan Fluor 40 × NA 1.3 lens was used. Excitation was from a 561 nm diode laser and emission was collected through a 605/75 nm bandpass filter.

3.2.11. Intracellular accumulation of nutlin-3a

MDCKII-pcDNA3.1 and MDCKII-BCRP cells were seeded into 6-well plates at a density of 0.5 × 10^5 cells/well in 2 ml of complete DMEM without G418. When cells were 80~90% confluent, nutlin-3a was added at increasing final concentrations of 0, 1, 5, 10, 20, and 50 μM. Cells were incubated at 37° C for 1 hour. Cells were then washed with ice cold phosphate buffered saline (PBS) twice, and scraped off in 1ml ice cold homogenization buffer (5 mM HCOONH4, pH 7.0). Cell pellets were then lysed on ice by sonicating for approximately 10 sec/well twice with a 15 sec interval using a 4710 series ultrasonic homogenizer (Cole-Parmer, Chicago, IL, USA). Intracellular nutlin-3a concentrations were measured using the LC-MS/MS method published previously [233]. Final intracellular nutlin-3a concentrations were normalized to total protein content as measured by BCA assay.

3.2.12. Bi-directional transport across MDCKII monolayer cells

MDCKII-pCDNA3.1 and MDCKII-BCRP cells were seeded at 1 × 10^6 cells per well in 0.4 μm, 12 mm Transwell® permeable inserts (Corning Incorporated, Corning, NY). Transport assays were performed when cells reached consistent trans-epithelial electrical resistance [234] values (between 200-300 μΩ cm^2), indicating that the cells had formed a confluent polarized monolayer. Prior to transport assays (30 minutes), the medium in the donor and receiver chambers was removed and replaced with transport buffer (HBSS/25 mM HEPES). Buffer containing nutlin-3a was added to either the apical or the basolateral side of the monolayer with or without the BCRP-specific inhibitor Ko143. Samples were removed for the determination of initial nutlin-3a concentrations, and then at 30, 60, 120, and 240 minutes. 50 μl aliquots were removed from the either the basolateral or apical compartments. The volume removed was replaced immediately with fresh transport buffer. Nutlin-3a samples were prepared and analyzed using LC-MS/MS method as described above.

The apparent permeability (Papp) is calculated using **Equation 3-1** [234]:

$$P_{\text{app}} = \frac{(V_r/C_0)}{(1/S)} (dC/dt)$$

**Eq. 3-1**
where \( V_r \) is the volume of buffer in the receiver chamber; \( C_0 \) is the initial drug concentration in the donor chamber; \( S \) is the surface area of monolayer; \( dC/dt \) is the linear slope of drug concentration in the receptor chamber over time.

The efflux ratio (RE) is calculated using **Equation 3-2**:

\[
RE = \frac{P_{app,B-A}}{P_{app,A-B}} 
\]

**Eq. 3-2**

where \( P_{app,B-A} \) is the apparent permeability of drug transport from basolateral to apical side and \( P_{app,A-B} \) is the apparent permeability of drug transport from apical to basolateral side.

The final efflux ratio (R) is calculated using **Equation 3-3**:

\[
R = \frac{\text{MDCK II-BCRP efflux ratio}}{\text{MDCK II-pcDNA 3.1 efflux ratio}}
\]

**Eq. 3-3**

### 3.2.13. ATPase assay

BCRP ATPase activity was determined using SB BCRP HAM PREDEASY™ ATPase kit and SB defBCRP HAM PREDEASY™ Ctrl kit following the manufacturer's instructions (XenoTech, Lenexa, KS). The assay contains two different tests that are performed simultaneously on the same plate. In the activation test, BCRP substrates stimulate baseline vanadate-sensitive ATPase activity. In the inhibition test, inhibitors or slowly transported compounds may inhibit the maximal vanadate sensitive ATPase activity. Nutlin-3a was tested in both the activation and inhibition reactions at increasing concentrations (0.14, 0.41, 1.23, 3.70, 11.11, 33.33, 100, and 150 µmol/L) for 10 minutes following the manufacturer's instructions. The assay was performed in triplicate.

### 3.2.14. Statistical analysis

All data expressed as mean ± standard deviation unless otherwise indicated. Data were analyzed for statistical significance using Student’s t test. Differences with \( p < 0.05 \) were considered statistically significant. Linear and nonlinear regressions were performed using Prism 5 (GraphPad Software, La Jolla, CA).

### 3.3. Results

#### 3.3.1. Nutlin-3a sensitizes BCRP expressing cells to mitoxantrone treatment

Saos-2-pcDNA3.1 and Saos-2-BCRP cells were incubated with nutlin-3a alone or in combination with the anti-cancer agent mitoxantrone, a BCRP substrate. \( IC_{50} \) values were determined by MTS assay. BCRP over-expression did not confer resistance to
nutlin-3a as a single agent. The IC$_{50}$ value of nutlin-3a was 45.8 (± 2.6) μM for Saos-2-BCRP and 43.5 (± 3.0) μM for Saos-2-pcDNA3.1 (p > 0.05) (Figure 3-1).

BCRP expression did confer resistance to mitoxantrone in the Saos-2 cell lines. The Saos-2-pcDNA3.1 cells exhibited sensitivity to mitoxantrone with an IC$_{50}$ of 2.0 (± 0.1) μM while the BCRP over-expressing Saos-2 cells had a markedly increased mitoxantrone IC$_{50}$ of 165.8 (± 21.9) μM (P < 0.001) (Figure 3-2). To determine the effect of nutlin-3a on potentially reversing this mitoxantrone resistance, Saos-2-BCRP cells were co-incubated with 20 μM or 50 μM nutlin-3a and increasing concentrations of mitoxantrone (0.01-300 μM) for 24 hours. A dramatic reduction in the mitoxantrone IC$_{50}$ from 165.8 (± 21.9) μM to 7.6 (± 0.5) μM (21.8-fold, p < 0.001) and 1.0 (± 0.07) μM (165.8-fold, p < 0.001) was observed in the Saos-2-BCRP cells after treatment with 20 and 50 μM nutlin-3a, respectively (Figure 3-2). In Saos-2-pcDNA3.1 cells, only a moderate reduction of mitoxantrone IC$_{50}$ was observed, from 2.0 (± 0.01) μM to 1.4 (± 0.1) μM (1.4-fold, p < 0.001) and 0.6 (± 0.4) μM (3.3-fold, p < 0.001) (Figure 3-2).

Using combination index analysis, we evaluated whether the combination of nutlin-3a and mitoxantrone was synergistic. Combination index (CI) values around 1 indicate that two drugs have an additive effect. A CI < 1 indicates synergy, and a CI > 1 indicates antagonism [226]. The fractional effect is the ratio of the effect (growth inhibition) caused by the two compounds in combination to that of one of the compounds alone. A fractional effect value of 0 indicates no inhibition and fractional effect value of 1 indicates 100% inhibition of cell viability. In contrast to Saos-2-pcDNA3.1 cells where synergism, additivity, and antagonism can be observed on different nutlin-3a: mitoxantrone ratios (Figure 3-3A), moderate (+) to strong (+++++) synergism [226] was observed in Saos-2-BCRP cells with combination index (CI) values between 0.132 and 0.798 at all nutlin-3a: mitoxantrone ratios tested (Figure 3-3B). These results indicate that the synergistic effect observed in Saos-2-BCRP cells induced by the nutlin-3a/mitoxantrone combination is dependent on the presence of BCRP.

### 3.3.2. Nutlin-3a inhibits BCRP-mediated transport of mitoxantrone

To determine whether the observed reduction in the mitoxantrone IC$_{50}$ of the Saos-2-BCRP cells co-treated with nutlin-3a was due to increased exposure to mitoxantrone, intracellular accumulation of mitoxantrone was measured by confocal imaging. Saos-2-pcDNA3.1 and Saos-2-BCRP cells were co-incubated with 1 μM mitoxantrone and increasing concentrations of nutlin-3a (0 to 50 μM) for 1 hour. In the absence of nutlin-3a, little mitoxantrone accumulation was observed in Saos-2-BCRP cells (Figure 3-4), indicating active efflux by BCRP, whereas under the same conditions, mitoxantrone was retained in the Saos-2-pcDNA3.1 cells. Treatment with nutlin-3a resulted in a dose-dependent increase in the intracellular mitoxantrone accumulation in the Saos-2-BCRP cells, suggesting a decrease in BCRP efflux function.
Figure 3-1. BCRP expression does not confer resistance to nutlin-3a treatment

Saos-2-BCRP and Saos-2-pcDNA3.1 cells were incubated with increasing concentrations of nutlin-3a (0.1-150 μM) for 24 hours. Cell viabilities were tested by MTS assay in triplicate. IC₅₀ values were calculated using ADAPT 5. Values are presented as mean ± SD. Data are representative of three independent experiments.
Figure 3-2. Co-treatment of cells with nutlin-3a and mitoxantrone strongly reverses BCRP mediated drug resistance to mitoxantrone

Saos-2-pcDNA 3.1 and Saos-2-BCRP cells were treated with increasing concentrations of mitoxantrone (0.01-300 μM) in combination with 20 and 50 μM nutlin-3a for 24 hours as described in the methods. Cell viabilities were determined by MTS assay in triplicate. ADAPT 5 was used to calculate IC$_{50}$. Values are presented as mean ± SD. *** P < 0.001. Data are representative of two independent experiments.
Figure 3-3. Synergistic effects of nutlin-3a in combination with mitoxantrone

The combination index (CI) values at non-fixed nutlin-3a/mitoxantrone concentration ratios were calculated using commercially available software Calcusyn 2.1. • CI values < 1.0 indicate synergism, ● CI values = 1.0 indicate additive effect, and □ CI values > 1.0 indicate antagonism. Fractional effect is defined as the fraction of cells affected by nutlin-3a and mitoxantrone combination. A fractional effect value of 0 indicates no inhibition and a fractional effect value of 1 indicates 100% inhibition of cell viability. Data are representative of two independent experiments.

A. For Saos-2-pcDNA 3.1 cells, at 20 μM nutlin-3a, additive effect and antagonism were observed. At 50 μM nutlin-3a, when nutlin-3a: mitoxantrone ratios were > 100:1, both additive and synergistic effects were observed; when nutlin-3a: mitoxantrone ratios were < 100:1, antagonism was observed.

B. For Saos-2-BCRP cells, moderate (++) to strong (++++) synergism was observed at all ratios tested.
Figure 3-4. Nutlin-3 treatment strongly increases the intracellular accumulation of mitoxantrone in Saos-2-BCRP cell lines

Nutlin-3 treatment strongly increases the intracellular accumulation of mitoxantrone in Saos-2-BCRP cell lines. Confocal imaging of mitoxantrone (red) in Saos-2-pcDNA3.1 and Saos-2-BCRP cells in the presence increasing concentrations of nutlin-3a for 60 minutes suggested dose dependent restoration of intracellular mitoxantrone accumulation. Scale bar, 30 μm. Data are representative of three independent experiments.
3.3.3. **Nutlin-3a inhibits BCRP-mediated transport of Hoechst 33342**

To determine whether nutlin-3a could also reduce the efflux of other BCRP substrates, the intracellular retention of another prototypical BCRP substrate, Hoechst 33342, was measured by flow cytometry in Saos-2 cells with and without BCRP expression. As BCRP over-expressing Saos-2 cells were incubated with increasing amounts of nutlin-3a, a dose-dependent decrease in the efflux of Hoechst 33342 was observed. Co-incubation of cells with nutlin-3a and Hoechst 33342 for 1 hour resulted in an almost complete inhibition of Hoechst 33342 efflux in Saos-2-BCRP cells (Figure 3-5A). This inhibition was comparable to that seen with the BCRP-specific inhibitor FTC (Figure 3-6). Co-incubation of cells with enantiomer nutlin-3b resulted in the same reduction in Hoechst 33342 efflux (Figure 3-7). Only events from viable cells were used for data analysis, and there was no difference in the viability between BCRP over-expressing and the vector control cells treated with either nutlin-3a or nutlin-3b (Figure 3-8). To determine if the reduced Hoechst 33342 efflux was dependent on p53 status, accumulation studies were also performed in p53 wild-type MDCKII cells. The results indicated that nutlin-3a also reverses the Hoechst 33342 efflux in a p53 wild-type cell line in a dose dependent manner (Figure 3-5B). Hence, nutlin-3a inhibition of BCRP-mediated transport of Hoechst 33342 is independent of cellular p53 status. In addition to the abrogation of efflux observed with flow cytometric methods, results from fluorescence imaging also demonstrated after treatment with nutlin-3a that Hoechst 33342 intracellular accumulation was dramatically restored in the Saos-2-BCRP cell line to levels comparable to Saos-2-pcDNA3.1 control cells (Figure 3-9). Treatment of cells with the enantiomer nutlin-3b resulted in the same reduction in Hoechst 33342 efflux, demonstrating that the two enantiomers have comparable effects on BCRP function (Figure 3-9).

3.3.4. **Nutlin-3a treatment does not alter BCRP expression or localization**

To determine whether increased accumulation and sensitivity to mitoxantrone were a result of nutlin-3a inducing an alteration in BCRP protein levels, Saos-2-pcDNA3.1 and Saos-2-BCRP cells were treated at the highest nutlin-3a dose level used for functional studies (50 μM; 1 hr) and BCRP protein levels were measured. As shown in Figure 3-10, both western blot (Figure 3-10A) and flow cytometric analysis (Figure 3-10B) demonstrated that total cellular protein levels of BCRP in Saos-2-BCRP cells were not altered in the presence of nutlin-3a. Since nutlin-3 has been shown to affect P-gp function, levels of P-gp protein were also assessed by western blot. Although Saos-2 cells do not express detectable levels of P-gp [232, 235], it is possible that nutlin-3a treatment may up-regulate P-gp expression. As expected, nutlin-3a treatment did not significantly alter the expression of P-gp in either Saos-2-pcDNA3.1 or -BCRP cells (data not shown). Additional studies using confocal microscopy confirmed no obvious alteration in the membrane translocation of BCRP protein (Figure 3-11).
Figure 3-5. Nutlin-3a dose-dependently inhibits BCRP efflux of Hoechst 33342, independent of p53 status

p53 null Saos-2-pcDNA3.1 and Saos-2-BCRP, and p53 wild-type MDCKII-pcDNA3.1 and MDCKII-BCRP cells were incubated with increasing concentrations of nutlin-3a or vehicle control in the presence of Hoechst 33342 for 60 minutes. Data are representative of at least three independent experiments.

A. Dose-dependent reduction of Hoechst 33342 efflux after nutlin-3a treatment assayed by flow cytometry in Saos-2 and MDCKII cells.

B. Quantitative assessment of nutlin-3a effects from representative flow cytometry experiments. Values are presented as mean ± SD.
Figure 3-6. Effect of FTC on the BCRP efflux of Hoechst 33342

Saos-2-pcDNA3.1 and Saos-2-BCRP cells were incubated with DMSO or FTC in the presence of Hoechst 33342 for 60 minutes. Data are representative of one experiment performed in triplicate. Error bar, ± SD. *p < 0.05, ***p < 0.0001.
Figure 3-7. Effect of nutlin-3b on efflux of Hoechst 33342

Saos-2-pcDNA3.1 and Saos-2-BCRP cells were incubated with DMSO or nutlin-3b in the presence of Hoechst 33342 for 60 minutes. Data are representative of one experiment performed in triplicate. Error bar, ± SD. ***p < 0.0001.
Figure 3-8. Viability ratio of Saos-2-BCRP/Saos-2-pcDNA3.1 during the Hoechst 33342 efflux study

Saos-2-pcDNA3.1 and Saos-2-BCRP cells were incubated with 50 μM nutlin-3a or nutlin-3b in the presence of Hoechst 33342 for 60 minutes. Data are representative of one experiment performed in duplicate. Error bar, ± SD. p > 0.05.
Figure 3-9. Nutlin-3 treatment increases the intracellular accumulation of Hoechst 33342 in Saos-2-BCRP cell lines

Wide-field fluorescence imaging of intracellular Hoechst 33342 (blue) in Saos-2-pcDNA3.1 and Saos-2-BCRP cells in the presence or absence of 50 μM nutlin-3a or nutlin-3b for 60 minutes. Treatment with nutlin-3a and nutlin-3b strongly restored the accumulation of Hoechst 33342 in Saos-2-BCRP cells. Data are representative of three independent experiments.
Figure 3-10. Nutlin-3a does not alter levels of BCRP protein expression

A. Western blot analysis of the BCRP protein expression following nutlin-3a treatment (N, 50 μM nutlin-3a; D, equal volume DMSO). BCRP protein was detected using the monoclonal antibody BXP-53.

B. Flow cytometric analysis of the cell-surface expression of BCRP in response to nutlin-3a treatment. Cells were stained with the PE conjugated mouse anti-BCRP antibody (MAB4155P) and subjected to analysis. DAPI was added to cells to indicate viability. Data are representative of two independent experiments.
Figure 3-11. Nutlin-3a does not alter cellular localization of BCRP in Saos-2 cells

Confocal images taken of Saos-2 cells expressing BCRP treated with vehicle (DMSO) or nutlin-3a (50 μM, 90 minutes). Red represents BCRP protein and blue represents nuclear staining. Scale bar, 30 μm. Data are representative of two independent experiments.
3.3.5. **Nutlin-3a is not a substrate for BCRP**

To test whether nutlin-3a is a BCRP substrate and thus inhibits BCRP function by competing with other substrates for transport, we examined the intracellular accumulation of nutlin-3a in MDCKII-pcDNA 3.1 and MDCKII-BCRP cells. Cells were incubated with 0, 1, 5, 10, 20, and 50 μM nutlin-3a at 37°C for 1 hour. No significant difference was observed in the amount of intracellular nutlin-3a between the two cell types at any concentration (up to 50 μM) (p > 0.05) (Figure 3-12). These data suggest that nutlin-3a is not a substrate of BCRP. In addition to the intracellular accumulation assay, bidirectional transport studies were conducted using BCRP over-expressing and pcDNA3.1 vector control MDCKII cells. Nutlin-3a concentrations from either the apical or basolateral compartment were measured by LC-MS/MS and concentration vs. time plots were generated (Figure 3-13). The calculated R (RE_{BCRP}/RE_{pcDNA3.1}) was 0.04, much less than the efflux ratio of 2, which is considered the cutoff for a drug to be a substrate [234]. Additionally, concentration vs. time plots of nutlin-3a in the presence and absence of the BCRP specific inhibitor Ko143 (5 μM) indicated that BCRP does not transport nutlin-3a (data not shown). Using the criteria outlined in the decision tree supported by the International Transporter Consortium, nutlin-3a is not a substrate of BCRP [47].

3.3.6. **Nutlin-3a inhibits the ATPase activity of BCRP**

To investigate inhibition as a mechanism for the nutlin-3a-induced reduction in efflux of BCRP substrates, BCRP ATPase activity in response to nutlin-3a treatment was measured using a previously described ATPase activity assay [93, 94]. BCRP transporter activity was determined by assaying both activation and inhibition in the presence of a known activator of the transporter (i.e., sulfasalazine). Increasing concentrations of nutlin-3a (0.1 μM to 150 μM) did not stimulate ATPase activities from the baseline measurements (data not shown). In the corresponding inhibition assay, however, higher concentrations of nutlin-3a demonstrated a strong capacity to inhibit ATPase activity (Figure 3-14).

3.4. **Discussion**

This is the first study demonstrating that nutlin-3a inhibits BCRP activity. Our data suggest that resistance to mitoxantrone can be strongly reversed by nutlin-3a in BCRP over-expressing cells. Nutlin-3a treatment resulted in a dose-dependent increase in the intracellular accumulation of BCRP substrates in BCRP over-expressing cells. To understand the mechanism behind these observations, a series of studies were performed that clearly demonstrated nutlin-3a inhibited BCRP efflux independent of p53, without altering BCRP protein expression or subcellular localization. Additionally, studies examining the intracellular accumulation of nutlin-3a along with bi-directional transport across MDCKII monolayer cells supported the conclusion that nutlin-3a is not a substrate of BCRP, but does act as an inhibitor through interference with BCRP ATPase activity.
MDCKII-pcDNA3.1 and MDCKII-BCRP cells were incubated with 0, 1, 5, 10, 20, and 50 μM nutlin-3a for 60 minutes. Intracellular nutlin-3a concentrations were determined by LC-MS/MS as previously described and normalized to total protein content. Data are presented as mean ± SD of triplicates.
Figure 3-13. Trans-epithelial transport of nutlin-3a (10 μM) in MDCKII-pcDNA3.1 and MDCKII-BCRP cells indicates nutlin-3a is not a substrate of BCRP

Nutlin-3a was administered to one compartment (basolateral or apical) at time 0. After 30, 60, 120, and 240 minutes, the concentrations of nutlin-3a appearing in the opposite compartment were measured by LC-MS/MS. Data are presented as mean ± SD of triplicates.

A. Translocation from the basolateral to the apical compartment. Data are representative of two independent experiments.

B. Translocation from the apical to the basolateral compartment. Data are representative of two independent experiments.
Figure 3-14. Nutlin-3a dose dependently inhibited BCRP ATPase activity

The relative vanadate-sensitive ATPase activity of Sf9 insect cell membranes over-expressing wild-type ABCG2 is represented as mol Pi/mg protein/min in the presence of increasing concentrations of nutlin-3a with a known BCRP substrate, sulfasalazine. Data are representative of three independent experiments.
Multi-drug resistance is a major obstacle in the success of cancer treatment. Among the ABC family of transporters, P-gp, MRP1, and BCRP are three major members associated with multidrug resistance [53]. BCRP, the most recently discovered among these three major transporters [53, 236], confers resistance to many anti-cancer drugs used clinically including mitoxantrone, methotrexate, doxorubicin, daunorubicin, topotecan, and SN38 [237, 238]. Utilization of an agent such as nutlin-3a, which inhibits BCRP, in combination with an anti-cancer agent that is a BCRP substrate (such as mitoxantrone or topotecan) may potentially increase the intracellular drug levels and lead to greater anti-tumor activity. In fact, when nutlin-3 was combined with the BCRP and P-gp substrate topotecan for 5 days, an 82-fold reduction in the tumor burden of retinoblastoma was reported [24]. It is important to note however that synergistic effects may differ depending on the cell type or co-administered drug, and antagonism may be observed if the schedule of administration were to change [239, 240].

Since nutlin-3a re-activates p53 in cells co-expressing MDM2 [17, 39], the question exists of whether the reversal of ABC transporter activity is dependent on the p53 pathway. Our data along with the previous study by Michaelis et al. clearly demonstrate that the inhibition of ABC transporter efflux by nutlin-3 occurs independently of cellular p53 status [228]. Also supporting this conclusion is the observation that nutlin-3b, the non-active enantiomer, demonstrated BCRP inhibition comparable to the active enantiomer nutlin-3a.

Michaelis et al. demonstrated that nutlin-3 stimulated P-gp ATPase activity in isolated membranes and exerted a negative effect on P-gp activity by acting as a competing substrate [228]. In contrast, our studies demonstrate by multiple approaches that nutlin-3a does not act as a competitive inhibitor of BCRP. First, the amount of intracellular nutlin-3a and the nutlin-3a IC50 were unaffected by the over-expression of BCRP. Second, the calculated efflux ratios from the bidirectional transport assay were < 2 in MDCKII cells. Lastly, nutlin-3a did not activate ATPase activity as measured by released inorganic phosphate (Pi) in a BCRP over-expression system. On the other hand, nutlin-3a dose dependently decreased the ATPase activity in the inhibition assay.

Previous studies have implicated BCRP translocation from the plasma membrane to the cytoplasm as a mechanism by which BCRP function can be regulated [241]. We demonstrated in our studies via western blot and flow cytometry that neither total BCRP protein levels nor the subcellular localization of BCRP changed during the time period that efflux studies were conducted. These findings are important because since BCRP is involved in drug disposition and many other physiological processes in the body, using a drug that alters BCRP expression and/or localization would likely have global effects. Specifically, BCRP is expressed at the blood brain barrier, placenta, gastrointestinal tract, kidney, liver and biliary tract, and BCRP activity is important for intestinal absorption, brain penetration, renal elimination and hepatobiliary excretion of substrates.

Along with BCRP, P-gp and MRP1 also play a critical role in pharmacokinetic interactions of anti-cancer agents, affecting the absorption, distribution, metabolism, and excretion (ADME) processes. Concomitant treatment of elacridar (GF120918), an
inhibitor of BCRP and P-gp, resulted in a 2.4 fold increase in bioavailability and systemic exposure of oral topotecan in adults with cancer [242]. Similarly, concomitant treatment with gefitinib, another inhibitor of BCRP and P-gp, increased bioavailability of oral irinotecan in mice [243] and in pediatric patients with refractory solid tumors [244]. Our lab recently demonstrated that gefitinib enhanced topotecan penetration in gliomas in mice [245]. As an inhibitor of multiple major efflux transporters including BCRP, P-gp and MRP1, nutlin-3a may have impact on pharmacokinetics, pharmacodynamics, and importantly the safety of many clinically used drugs. Therefore, it is crucial that transporter related drug-drug interactions be carefully addressed in future preclinical studies.

In conclusion, this is the first study demonstrating that nutlin-3a inhibits BCRP activity. Our data show that nutlin-3a dose dependently inhibits BCRP-mediated transport of multiple BCRP substrates and synergistically reverses the drug resistance to anticancer agent mitoxantrone. The likely mechanism of this effect is the inhibition of BCRP ATPase activity, as we have clearly demonstrated through multiple lines of investigation that nutlin-3a is not a substrate of BCRP. Thus, using nutlin-3a in combination with anti-cancer agents that are BCRP or other ABC transporter substrates would require additional studies to identify potentially significant drug-drug interactions due to the critical role of these transporters in drug ADME.
CHAPTER 4. WHOLE-BODY PHYSIOLOGICALLY BASED
PHARMACOKINETIC MODEL FOR NUTLIN-3A IN MICE AFTER
INTRAVENOUS AND ORAL ADMINISTRATION*

4.1. Introduction

Nutlin-3a (2-piperazinone, 4-[[4S,5R]-4,5-bis(4-chlorophenyl)-4,5-dihydro-2-[4
methoxy-2-(1-methylethoxy)phenyl]-1H-imidazol-1-yl]carbonyl]-) is currently
undergoing preclinical investigation as a p53 reactivation agent. While many cancers and
tumor types express mutated forms of p53 [246], a subset of cancers, and particularly
pediatric tumors, retain wild-type p53 [211]. In these cases, cancer cells frequently
employ other mechanisms to abrogate p53 function. One such mechanism is
over-expression or amplification of the murine double minute (MDM2) protein. This
molecule binds directly to p53 to accelerate its turnover and inhibits transcription of
downstream targets, including cell cycle and apoptotic genes [8, 218]. Disruption of the
MDM2–p53 interaction is proposed as a novel strategy for treatment of cancers that do
not have p53 alterations [21, 247].

Nutlins are a class of small molecules that target the p53-binding pocket of
MDM2 [17, 248]. Treatment of multiple types of cancer cells including leukemias [26,
41], neuroblastoma [221], rhabdomyosarcoma [249] and retinoblastoma [250] with
nutlin-3a induces p53-dependent cell cycle arrest and cell death, whereas in normal cells,
nutlin-3a exposure leads to cell cycle arrest without cell death [22]. Nutlin-3a has
antitumor activity in a preclinical xenograft model of neuroblastoma [43] and was tested
in several other preclinical models of malignancies [17, 40].

To date, the pharmacokinetics of nutlin-3a has not been reported. An
understanding of the systemic disposition of nutlin-3a, as well as the distribution to target
tissue or tumor sites, will provide a rational basis for the selection of dosage regimens for
preclinical models. Additionally, since the in vitro tumor cell line sensitivities to
nutlin-3a have been determined, pharmacokinetic modeling can be used to determine the
dose and schedule necessary to achieve appropriate unbound nutlin-3a concentrations at
the tumor site. One approach to analyze these data is the use of whole-body
physiologically-based pharmacokinetic model (PBPK) models, which are based on
anatomical compartments and blood flow.

Thus, we performed pharmacokinetic studies to develop a PBPK model
describing the disposition of nutlin-3a in plasma and tissues, including adipose, adrenal
gland, bone marrow, brain, liver, lung, intestine, muscle, retina, spleen, and vitreous
fluid. The PBPK model was used to perform simulations, which in combination with in
vitro cell sensitivity data provided rationale for choosing dosing regimens for mouse

* Reprinted with permission. Zhang, F., et al., Whole-body physiologically based
pharmacokinetic model for nutlin-3a in mice after intravenous and oral
models of common childhood cancers, including retinoblastoma, neuroblastoma, rhabdomyosarcoma, and acute lymphoblastic leukemia (ALL).

4.2. Materials and Methods

4.2.1. Animals

Adult C57BL/6 mice were purchased from Charles River (Bar Harbor, Maine, USA). Mice were housed in a temperature-controlled room on a normal 12-h light/dark cycle, with free access to water and standard laboratory food. All procedures were approved by the St. Jude Institutional Animal Care and Use Committee and conducted in accordance with the NIH guidelines for the care and use of laboratory animals. The animal facility is accredited by the American Association for Accreditation of Laboratory Animal Care.

4.2.2. Chemicals

cis-Nutlin-3a (98% purity, Lot No.: 08252008) was synthesized and supplied by the Department of Chemical Biology and Therapeutics at St. Jude Children’s Research Hospital (Memphis, TN, USA). The oral formulation [17] used in the pharmacokinetic studies was nutlin-3a suspended in 2% Klucel (Conservation Resources International, LLC, Springfield, VA, USA) and 0.5% Tween-80 (Sigma-Aldrich, St. Louis, MO, USA), and the IV formulation used was nutlin-3a in 4% ethanol, 35% propylene glycol (Fisher Scientific, Pittsburgh, PA, USA), 10% PEG-400 (Sigma-Aldrich), and 51% PBS (Mediatech INC, Manassas, VA, USA). The internal standard ketoconazole was purchased from Sigma-Aldrich. Blank murine plasma was obtained from Hilltop Lab Animals, Inc. (Scottsdale, PA, USA). All other reagents were of analytical grade or higher.

4.2.3. Blood to plasma ratio

Pooled whole blood from healthy male C57BL/6 mice was collected into heparin tubes. Nutlin-3a was spiked into aliquots of whole blood at final concentrations of 0.1, 1, 10, and 100 μM in triplicate. Samples were mixed and incubated at 37°C for 30 minutes with additional mixing every 5 minutes. After incubation, 50 μl of whole blood was removed and immediately placed on dry ice. The remainder of the sample was centrifuged at 16,000 rpm for 2 min and a 50 μl plasma sample was removed and immediately placed on dry ice. All samples were stored at -80°C until analysis. The blood to plasma concentration ratio was calculated using the using \textbf{Equation 4-1}:

\[
\text{Blood to plasma ratio} = \frac{c_{WB}}{c_p} \quad \text{Eq. 4-1}
\]
where $C_P$ is the concentration in plasma, $C_{WB}$ is the concentration in whole blood.

4.2.4. Nutlin-3a protein binding studies

Equilibrium dialysis was performed in a 96-well dialysis plate with a 5-KDa cut-off membrane (Harvard Apparatus, Holliston, MA, USA). For mouse plasma and cell culture media protein binding, 200 μl of PBS buffer was added into the wells on one side of the membrane and an equivalent volume of male C57BL/6 plasma or cell culture media (RPMI with 10% FBS and 2 mM L-glutamine) containing varying concentration of nutlin-3a was added into the wells on the opposite side. The plate was sealed and fixed onto a dual plate rotator (Harvard Apparatus) at 37ºC in a humidified incubator containing 5% CO₂. Equilibrium dialysis was performed at 0.2, 20, and 100 μM nutlin-3a in triplicate for 24 h. For vitreous protein binding, 150 μl rodent vitreous containing 0.5 μM nutlin-3a was added to the sample side and equal volume of PBS buffer was added to the buffer side. Equilibrium dialysis was performed in triplicate for 24 h. The samples were analyzed using the analytical method described below. The bound concentration was considered equal to the plasma side and the free concentration equal to the PBS side [251, 252]. Binding parameters were estimated with nonlinear regression using Equation 4-2 [253]:

$$C_{p,b} = \frac{B_{\text{max}}K_A C_{p,f}}{1 + K_A C_{p,f}}$$

Eq. 4-2

where $C_{p,b}$ is the bound plasma concentration, $B_{\text{max}}$ is the quantity of plasma protein binding sites, $K_A$ is the binding association constant, and $C_{p,f}$ is the unbound plasma concentration.

4.2.5. Drug administration and sample collection

Two pharmacokinetic studies were conducted. For the first pharmacokinetic study, 145 adult C57BL/6 mice (128 male and 17 female) were divided into three groups: two oral dosage groups (100 and 200 mg/kg) and one IV dosage group (10 mg/kg). Nutlin-3a was administered given as a single bolus dose by oral gavage or by i.v. tail vein injection. Each dosing group (n = 5 mice) and vehicle controls had 9 collection time points (0.5, 1, 2, 4, 8, 12, 24, 36, and 48 h). At each time point, blood was collected under isoflurane anesthesia via cardiac puncture. Whole blood samples were centrifuged immediately at 12,000 g at 4 ºC for 5 minutes to separate plasma. Simultaneously, tissue samples including brain, vitreous, retina, liver, spleen, and bone marrow were dissected. Each sample was put on dry ice immediately after collection and stored at -80ºC until analysis.

In the second pharmacokinetic study, 210 adult male C57BL/6 mice were used. Two oral dosages (50 and 100 mg/kg) and two i.v. dosages (10 and 20 mg/kg) were administered. Each dosing group (n = 5 for 10 mg/kg IV and 100 mg/kg oral dosages;
n = 10 for 20 mg/kg IV and 50 mg/kg oral dosages) and vehicle control had 7 collection
time points (0.5, 1, 2, 4, 8, 12, and 24 h for the IV dosing; 0.5, 1, 2, 4, 8, 12, and 16 h for
oral dosing). Serial plasma samples were collected from all mice. Tissue samples,
including brain, lung, liver, spleen, kidney, adrenal gland, muscle, fat, intestine from 3
mice per time point from the 20 mg/kg IV group, were collected. Each sample was put on
dry ice immediately after the sample collection and stored at -80ºC until analysis.

4.2.6. Quantitative analysis of nutlin-3a

Nutlin-3a mouse plasma and tissue samples were analyzed based on our
previously published liquid chromatography electrospray ionization tandem mass
spectrometry analytical method [233]. The lower limit of quantification (LLOQ) for
nutlin-3a in plasma after protein precipitation was 10 ng/ml and the LLOQ for nutlin-3a
after liquid-liquid extraction was 0.25 ng/mL. Within-day and between-day precisions for
protein precipitation and liquid-liquid extraction were ≤ 5% and accuracies ranged from
91.6 to 104.8%. For each sample type (cerebellum, brain, vitreous, retina, lung, heart,
liver, gall bladder, spleen, kidney, adrenal gland, muscle, fat, bone marrow, intestine,
whole blood, and PBS), standard curves and controls were generated using the
corresponding untreated tissue or PBS to eliminate any matrix effect. For larger tissues,
sections were cut, weighed, and stored on ice for further processing. 10 μL ice cold
homogenization buffer (5 mM HCOONH4, pH = 7) were added per mg of tissue. For
smaller tissue samples including vitreous, retina, adrenal gland, and gall bladder, the
amount of homogenization buffer used was increased to a minimum volume of 70~100
μL. Tissue samples were then sonicated on ice for 15 seconds, with 5 second intervals.
The number of total sonications varied depending on the tissue types. Homogenated
tissues and whole blood samples were extracted and analyzed using protein precipitation
and the PBS samples were extracted using the liquid-liquid extraction method as
described previously [233].

4.2.7. Whole body PBPK model development

We developed a whole body PBPK model for nutlin-3a based on *in vitro* blood
cell partitioning, plasma protein binding, and pooled concentration-time data from all
plasma and tissue samples collected from both pharmacokinetic studies. This PBPK
model consisted of a series of mass balance differential equations describing the
concentration of nutlin-3a in various tissues, which were connected by blood flow.
Physiological values for mouse organ size and blood flow are presented in Table 4-1. A
schematic representation of the model is shown in Figure 4-1.

Plasma concentrations were converted to whole blood concentrations based on the
*in vitro* blood partitioning experiment. Unbound plasma concentrations were described
with Equation 4-3 [253]:

72
Table 4-1. List of physiological parameters

<table>
<thead>
<tr>
<th>Tissue</th>
<th>Symbol</th>
<th>Mass (% body weight)</th>
<th>(Q_B) (ml/h)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood</td>
<td>BLO</td>
<td>4.9</td>
<td>839</td>
<td>[254]</td>
</tr>
<tr>
<td>Adipose</td>
<td>ADI</td>
<td>6.8</td>
<td>58.7</td>
<td>[254]</td>
</tr>
<tr>
<td>Adrenal gland</td>
<td>ADR</td>
<td>0.048</td>
<td>2.52</td>
<td>[254]</td>
</tr>
<tr>
<td>Bone marrow</td>
<td>MRW</td>
<td>5.8</td>
<td>92.3</td>
<td>[254]</td>
</tr>
<tr>
<td>Brain</td>
<td>BRA</td>
<td>1.65</td>
<td>27.7</td>
<td>[254]</td>
</tr>
<tr>
<td>Intestines</td>
<td>INT</td>
<td>3.62</td>
<td>109</td>
<td>[254]</td>
</tr>
<tr>
<td>Liver</td>
<td>LIV</td>
<td>5.49</td>
<td>16.8</td>
<td>[254]</td>
</tr>
<tr>
<td>Lung</td>
<td>LUN</td>
<td>0.73</td>
<td>839</td>
<td>[254]</td>
</tr>
<tr>
<td>Muscle</td>
<td>MUS</td>
<td>38.4</td>
<td>133</td>
<td>[254]</td>
</tr>
<tr>
<td>Retina</td>
<td>RET</td>
<td>0.04</td>
<td>3.16</td>
<td>Experimental; [255]</td>
</tr>
<tr>
<td>Spleen</td>
<td>SPL</td>
<td>0.35</td>
<td>9.48</td>
<td>[254]</td>
</tr>
<tr>
<td>Vitreous fluid</td>
<td>VIT</td>
<td>0.035</td>
<td>0*</td>
<td>Experimental</td>
</tr>
<tr>
<td>Remainder</td>
<td>RES</td>
<td>29.9</td>
<td>256.9</td>
<td>-</td>
</tr>
</tbody>
</table>

* Vitreous assumed to have no direct blood flow.
Figure 4-1. Schematic diagram of PBPK model for nutlin-3a in mice

C,b = bound drug concentration, C,f = free drug concentration. Arrows connecting compartments represent blood flows from literature values.
\[ C_{p,u} = \frac{-(1+B_{\text{max}} - K_A \cdot C_p) + \left((1+B_{\text{max}} - K_A \cdot C_p)^2 + 4 \cdot K_A \cdot C_p\right)^{1/2}}{2 \cdot K_A} \]  

Eq. 4-3

where \( C_{p,u} \) is the unbound plasma concentration, \( C_p \) is the total plasma concentration, and \( B_{\text{max}} \) and \( K_A \) were determined from \textit{in vitro} plasma protein binding studies. The unbound fraction \( (f_{ub}) \) was calculated by \textbf{Equation 4-4}:

\[ f_{ub} = \frac{C_{p,u}}{C_p} \]  

Eq. 4-4

Most organs fit well to a perfusion-limited model, and thus were described by \textbf{Equation 4-5}:

\[ V_i \cdot \frac{dA_i}{dt} = Q_i \cdot \left(C_{\text{ART}} - \frac{C_i}{K_i}\right) \]  

Eq. 4-5

where \( V_i \) is the volume of organ, \( A_i \) is the amount of drug in the organ, \( C_i \) is the concentration in the organ, \( K_i \) is the partition coefficient, and \( C_{\text{ART}} \) is the concentration of arterial plasma.

Liver blood flow \( (Q_{\text{LIV}}) \) was the sum of the blood flow to the hepatic artery, spleen, and liver, and the concentration of blood entering the liver \( (C_{\text{BLO}, \text{LIV}}) \) was based on the arterial concentration and the venous outflow of the portal circulation (\textbf{Equation 4-6}). The liver contained an elimination term \( (k_e) \) for metabolism, based on experiments showing that nutlin-3a is metabolized by mouse liver microsomes (K. Guy, unpublished):

\[ V_{\text{LIV}} \cdot \frac{dA_{\text{LIV}}}{dt} = Q_{\text{LIV}} \cdot \left(C_{\text{BLO}, \text{LIV}} - \frac{C_{\text{LIV}}}{K_{\text{LIV}}}\right) - k_e \cdot C_{\text{ART}} \]  

Eq. 4-6

The intestine was modeled with a separate lumen and tissue compartment. Absorption from the lumen was assumed to be linear based on an absorption rate constant \( (k_a) \) (\textbf{Equation 4-7}):

\[ V_{\text{INT}} \cdot \frac{dA_{\text{INT}}}{dt} = k_a \cdot A_{\text{INT}} \]  

Eq. 4-7

The eye was fit to a two-compartment model consisting of the retina and vitreous. Input into the vitreous was by diffusion from the retina. The following equations were used for the retina and vitreous (\textbf{Equation 4-8} and \textbf{Equation 4-9}):

\[ V_{\text{RET}} \cdot \frac{dA_{\text{RET}}}{dt} = Q_{\text{RET}} \cdot \left(C_{\text{ART}} - \frac{C_{\text{RET}}}{K_{\text{RET}}}\right) - PA_{\text{VIT}} \cdot \left(C_{\text{RET}} - \frac{C_{\text{VIT}}}{K_{\text{VIT}}}\right) \]  

Eq. 4-8

\[ V_{\text{VIT}} \cdot \frac{dA_{\text{VIT}}}{dt} = PA_{\text{VIT}} \cdot \left(C_{\text{RET}} - \frac{C_{\text{VIT}}}{K_{\text{VIT}}}\right) \]  

Eq. 4-9

where \( PA_{\text{VIT}} \) is the permeability-surface area product.
All tissues that were not sampled were lumped together in a residual compartment. Modeling this compartment as perfusion-limited did not adequately describe the multi-exponential profile of nutlin-3a. Therefore, the residual compartment was modeled as diffusion-limited, with the vascular space assumed to be 5% of the residual volume. The equations for the residual vascular space and tissue are described using \textbf{Equation 4-10} and \textbf{Equation 4-11}:

\begin{equation}
V_{RES,BLO} \cdot \frac{dA_{RET}}{dt} = Q_{RES} \cdot \left( C_{ART} - C_{RES,BLO} \right) - PA_{RES} \cdot \left( C_{RES} - \frac{C_{RES}}{K_{RES}} \right) \quad \text{Eq. 4-10}
\end{equation}

\begin{equation}
V_{RES} \cdot \frac{dA_{RES}}{dt} = PA_{RES} \cdot \left( C_{RES} - \frac{C_{RES}}{K_{RES}} \right) \quad \text{Eq. 4-11}
\end{equation}

The input into the venous pool of blood was modeled as the sum of the output from all tissues except the lung. The volume of the venous pool was fixed to 75% of the total blood volume. The lungs received all venous input and the arterial input was the output from the lungs (\textbf{Equation 4-12}):

\begin{equation}
V_{LUN} \cdot \frac{dA_{LUN}}{dt} = Q_{BLO} \cdot \left( C_{VEN} - \frac{C_{LUN}}{K_{LUN}} \right) \quad \text{Eq. 4-12}
\end{equation}

The arterial concentrations were based on the output from the lungs, and contained an additional saturable elimination term (\textbf{Equation 4-13}):

\begin{equation}
V_{ART} \cdot \frac{dA_{ART}}{dt} = Q_{BLO} \cdot \left( \frac{C_{LUN}}{K_{LUN}} - C_{ART} \right) - \frac{V_{max} \cdot C_{ART}}{K_{m} + C_{ART}} \quad \text{Eq. 4-13}
\end{equation}

Elimination terms in both the blood compartment and liver compartment were necessary for a good model fit to the data from both oral and intravenous administration.

\subsection*{4.2.8. Simulations}

After development of the PBPK model, tissue concentrations were simulated with NONMEM after multiple oral and intravenous doses at 50, 100, 200, and 400 mg/kg given both once daily and twice daily. The AUC\textsubscript{0-24} was calculated with the log-linear trapezoidal method applied to the simulated data. Bioavailability was estimated using the ratio of AUC\textsubscript{0-24, IV}/AUC\textsubscript{0-24, PO} with the simulated steady state data.

\subsection*{4.3. Results}

\subsection*{4.3.1. Blood to plasma partitioning and plasma protein binding of nutlin-3a}

Blood to plasma partitioning showed an average blood/plasma concentration ratio of 0.70, indicating that 30% of nutlin-3a partitions to blood cells (\textbf{Figure 4-2A}). Binding of nutlin-3a to mouse plasma proteins was nonlinear, with $f_{ub}$ ranging from 0.007 at 0.1
Figure 4-2. Analysis of nutlin-3a characteristics in murine blood

A. Nutlin-3a blood cell partitioning. Nutlin-3a was spiked into murine whole blood at various concentrations, and incubated 30 min at 37ºC. In one aliquot, nutlin-3a was measured in whole blood, and in another aliquot nutlin-3a was measured in the plasma.

B. Nutlin-3a plasma protein binding. Nutlin-3a was spiked into murine plasma at various concentrations and incubated for 30 min at 37ºC. Plasma protein binding was determined by equilibrium microdialysis and is expressed as the percent of the total nutlin-3a plasma concentration that is unbound. Bars represent the mean and error bars represent the standard deviation.
μM to 0.118 at 300 μM (Figure 4-2B). Nonlinear regression of unbound versus bound plasma concentrations using the Langmuir equations resulted in a Bmax of 286 and a KA of 0.085 (Figure 4-3).

4.3.2. Nutlin-3a pharmacokinetics in mice

Plasma and tissue concentrations of nutlin-3a were measured from 0 to 48 h in mice following a single i.v. dose of 10 or 20 mg/kg or a single oral dose of 50, 100, or 200 mg/kg. After oral administration, nutlin-3a tissue concentrations rose rapidly to reach a maximum value at approximately 2 h. Nutlin-3a concentrations in the intestine, liver, and spleen were higher than those in the plasma, concentrations in adipose, adrenal gland, lung, muscle, and retina were similar to plasma concentrations, and concentrations in the brain, bone marrow, and vitreous were significantly lower than in the plasma (Figure 4-4).

Rapid elimination was observed in the 10 mg/kg IV dosage group. At higher dosages, slower elimination was observed at higher concentrations, indicating saturable elimination of nutlin-3a. After 24 h, all data were below the limit of quantitation of the assay. Models with linear elimination, Michaelis-Menten elimination, and combined linear and Michaelis-Menten elimination were fit to the data. Ultimately, the combination of both linear and Michaelis-Menten elimination had the best fit to the data. The concentration-time data of nutlin-3a in all modeled tissues are plotted against the model predicted concentrations in Figure 4-5 (data not shown for oral 50 mg/kg dosage group, since only plasma was collected). The estimated pharmacokinetic parameters are listed in Table 4-2.

Using the final model, we simulated nutlin-3a plasma concentrations after multiple doses on a once-daily and twice-daily schedule (Figure 4-6). Little to no accumulation was predicted to occur on a once daily schedule with IV or oral dosages up to 400 mg/kg. For twice daily dosing, steady state was predicted to occur within three doses, but accumulation would remain minimal. The predicted accumulation (Cmin single dose/Cmin at steady state) was dose-dependent and at 200 mg/kg is 1.5-fold for IV administration and 1.3-fold for oral administration. The AUC0-24 at steady state increased in an approximately dose-proportional manner and the AUC0-24 was approximately twice as high with twice-daily dosing versus once-daily dosing (Figure 4-7).

Predicted bioavailability was dose- and schedule -dependent and ranged from 75% at 25 mg/kg once-daily to 91% at 400 mg/kg once-daily. Bioavailability was predicted to be near 100% when given twice daily at dosages of 50 mg/kg or higher.
Figure 4-3. Nutlin-3a binding to murine plasma proteins

Bound and unbound nutlin-3a plasma concentrations were determined with equilibrium dialysis and nonlinear regression was performed using the Langmuir equation.
Figure 4-4. Comparison of actual plasma and tissue concentrations of nutlin-3a

The median plasma concentrations are shown in each box as the dashed line and the symbols represent the median tissue concentrations.
Figure 4-5. Concentration-time plots of nutlin-3a in tissues

Symbols are data points from individual mice and the lines represent the model-predicted concentrations. Data from the 50 mg/kg oral group is not shown. Data below the lower limit of quantitation is not shown.
### Table 4-2. Estimated PBPK model parameters

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
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</thead>
<tbody>
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</tr>
<tr>
<td>$K_e$</td>
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</tr>
<tr>
<td>$V_{max}$</td>
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</tr>
<tr>
<td>$K_{VIT}$</td>
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</tr>
<tr>
<td>PA_{VIT}</td>
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</tr>
<tr>
<td>$K_{RES}$</td>
<td>4.8</td>
</tr>
<tr>
<td>PA_{RES}</td>
<td>0.0048</td>
</tr>
<tr>
<td>IIV $k_a$</td>
<td>31.2%</td>
</tr>
<tr>
<td>IIV $k_e$</td>
<td>6.4%</td>
</tr>
<tr>
<td>IIV $V_{max}$</td>
<td>40.6%</td>
</tr>
<tr>
<td>Residual error</td>
<td>35.6%</td>
</tr>
</tbody>
</table>
Figure 4-6. Simulated concentration-time plot of plasma nutlin-3a after multiple oral doses with once-daily (QD) and twice-daily (BID) dosing

Simulations were based on the final PBPK model.
Figure 4-7. Plasma area under the concentration-time curve for 24 h at steady state (AUC0-24) versus nutlin-3a dosage when administered once daily (QD) and twice daily (BID).

AUCs were calculated from simulated concentration-time curves based on the final PBPK model.
4.3.3. Application of PBPK model to the design of nutlin-3a dosing regimens in mice

The nutlin-3a PBPK model was used to choose dosing regimens of nutlin-3a to target (1) the retina and vitreous for models of retinoblastoma, (2) the adrenal gland for models of neuroblastoma, (3) the muscle for models of rhabdomyosarcoma, and (4) the plasma, spleen, and bone marrow for models of leukemia. The fraction of unbound nutlin-3a in tissues was assumed to be the same as the plasma unbound fraction, except for vitreous fluid that had a measured unbound fraction of 14.4%. Nutlin-3a binding to cell culture media was measured and shown to be nonlinear over the range of nutlin-3a concentrations used in in vitro cell cytotoxicity assays (Figure 4-8). The media protein binding value was used to convert the published nutlin-3a IC$_{50}$ values of various cell lines [26, 249, 256] to the unbound IC$_{50}$ (Table 4-3).

The simulated unbound retina and vitreous nutlin-3a concentrations were compared to the in vitro unbound IC$_{50}$ for the Weri1 retinoblastoma cell line in order to choose optimal dosing regimens for mouse models of retinoblastoma (Figure 4-9). Oral dosing of nutlin-3a twice daily was predicted to achieve unbound concentrations in the retina that were consistently above the unbound IC$_{50}$ at dosages of 200 or 400 mg/kg (Table 4-4). However, even at 400 mg/kg twice daily, unbound concentrations in the vitreous were predicted to be above the unbound IC$_{50}$ for only 17% of the time, and at lower dosages the concentration of unbound nutlin-3a never reached the unbound IC$_{50}$ level. Simulated concentration-time plots of unbound nutlin-3a after various dosing regimens are also shown in the adrenal gland for neuroblastoma (Figure 4-10), muscle for rhabdomyosarcoma (Figure 4-11) and plasma, bone marrow, and spleen for leukemia (Figure 4-12). The percent times above the unbound IC$_{50}$ are also listed in (Table 4-4).

4.4. Discussion

Nutlin-3a is undergoing preclinical studies examining its potential efficacy for the treatment of several childhood malignancies. Nutlin-3a interrupts the p53-MDM2 protein-protein interaction, which may lead to apoptosis or cell cycle arrest. Treatment of cells with nutlin-3a leads to reversal of multi-drug resistance [228], reduced cell migration [229], reduced angiogenesis [214, 229], radiosensitization of hypoxic cancer cells [29], and inhibition of tumor adaptation to hypoxia [257]. In this study, we developed a mouse PBPK model of nutlin-3a in plasma and multiple tissues of therapeutic interest. This is the first study to provide comprehensive pharmacokinetic data of nutlin-3a in any species. The design of our study included both oral and intravenous dosing at several dosage levels. This permitted the development of a robust model that accurately describes the disposition of nutlin-3a over a wide range of concentrations. The PBPK model was used to design rationale dosing regimens for preclinical models of several malignancies based upon achieving adequate cytotoxic nutlin-3a concentrations within the target organ. This approach is superior to optimizing dosing based solely on plasma concentrations since drug penetration to different organs can vary widely.
Figure 4-8. Nutlin-3a binding to cell culture media

Bound and unbound nutlin-3a plasma concentrations were determined with equilibrium dialysis.

A. The unbound fraction is shown with increasing total nutlin-3a concentrations. Bars represent the mean and error bars represent the standard deviation for one experiment performed in triplicate.

B. Nonlinear regression was used to fit a binding model (Langmuir equation) to the data.

Table 4-3. Nutlin-3a IC\textsubscript{50} for cell survival in different cell types

<table>
<thead>
<tr>
<th>Cell type</th>
<th>IC\textsubscript{50} (\textgreek{M})</th>
<th>Unbound IC\textsubscript{50} (\textgreek{M})</th>
<th>Exposure time (h)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Weri1 retinoblastoma</td>
<td>1.1</td>
<td>0.21</td>
<td>72</td>
<td>Unpublished\textsuperscript{a}</td>
</tr>
<tr>
<td>IMR-32 neuroblastoma</td>
<td>3.02</td>
<td>0.77</td>
<td>72</td>
<td>[221]</td>
</tr>
<tr>
<td>RMS-YM rhabodmyosarcoma</td>
<td>1.25\textsuperscript{b}</td>
<td>0.26</td>
<td>48</td>
<td>[249]</td>
</tr>
<tr>
<td>Primary MDM2-overexpressing ALL</td>
<td>1.0\textsuperscript{b}</td>
<td>0.20</td>
<td>44</td>
<td>[26]</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Mike Dyer, St. Jude Children’s Research Hospital, Memphis, TN.

\textsuperscript{b} Nutlin-3a IC\textsubscript{50} assumed to be half of racemic nutlin-3 IC\textsubscript{50}.
Figure 4-9. Simulated concentration-time plot of unbound nutlin-3a in the retina and vitreous after multiple oral doses given once daily (QD) or twice daily (BID)

The horizontal lines represent the unbound IC$_{50}$ of nutlin-3a for Weri1 cells.

Table 4-4. Percent time unbound tissue concentration is above unbound IC$_{50}$

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Tissue</th>
<th>Time above IC$_{50}$ (%)</th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>QD 100</td>
<td>QD 200</td>
<td>QD 400</td>
<td>BID 100</td>
<td>BID 200</td>
<td>BID 400</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Weri1</td>
<td>Retina</td>
<td>38</td>
<td>53</td>
<td>70</td>
<td>83</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Vitreous</td>
<td>0</td>
<td>0</td>
<td>5</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>17</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>IMR32</td>
<td>Adrenal</td>
<td>27</td>
<td>39</td>
<td>54</td>
<td>58</td>
<td>85</td>
<td>100</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RMS-YM</td>
<td>Muscle</td>
<td>43</td>
<td>58</td>
<td>76</td>
<td>92</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ALL</td>
<td>Plasma</td>
<td>43</td>
<td>57</td>
<td>75</td>
<td>90</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Spleen</td>
<td>46</td>
<td>61</td>
<td>80</td>
<td>90</td>
<td>100</td>
<td>100</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Marrow</td>
<td>0</td>
<td>23</td>
<td>35</td>
<td>12</td>
<td>48</td>
<td>77</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

QD: once daily dosing. BID: twice daily dosing.
Figure 4-10. Simulated concentration-time plot of unbound nutlin-3a in the adrenal gland after multiple oral doses given once daily (QD) or twice daily (BID)

The horizontal lines represent the unbound IC$_{50}$ of nutlin-3a for IMR-32 p53-wt neuroblastoma cells.

Figure 4-11. Simulated concentration-time plot of unbound nutlin-3a in the muscle after multiple oral doses given once daily (QD) or twice daily (BID)

The horizontal lines represent the unbound IC$_{50}$ of nutlin-3a for RMS-YM rhabdomyosarcoma cells.
Figure 4-12. Simulated concentration-time plot of unbound nutlin-3a in the plasma, bone marrow, and spleen after multiple oral doses given once daily (QD) or twice daily (BID).

The horizontal lines represent the unbound IC$_{50}$ of nutlin-3a for MDM2-overexpressing primary acute lymphoblastic leukemia cells.
Our model describes the key pharmacokinetic properties of nutlin-3a in plasma: rapid absorption, high bioavailability, and saturable elimination that is very rapid at concentrations below 10 μM. Standard non-compartmental calculations of bioavailability of nutlin-3a were greater than 100% due to saturable elimination and different ranges of intravenous (10-20 mg/kg) and oral (50-100 mg/kg) dosages. Using the model to simulate the concentration-time profiles after the same dosage administered both intravenous and oral, we were able to estimate nutlin-3a oral bioavailability. Although we performed simulations after multiple doses, the model was based on data from only single doses of nutlin-3a, and therefore should be interpreted with caution.

Nutlin-3a is a substrate for the ATP binding cassette (ABC) transporter P-glycoprotein (P-gp), but at higher concentrations can also inhibit P-gp efflux activity [228]. The inhibition of P-gp function may explain why nutlin-3a is capable of rapid absorption and high bioavailability despite being a P-gp substrate. It is also possible that inhibition of P-gp function underlies saturable nutlin-3a elimination, since P-gp can excrete drugs into both the bile and urine [47]. It is unknown whether nutlin-3a may also inhibit its own metabolism at higher concentrations.

The partition coefficients showed a greater than 1000-fold difference between the tissues with the lowest and highest penetration. The liver and intestine showed the highest penetration. High penetration to the liver could be due to uptake transporters expressed at the hepatocyte membranes, which cause intracellular accumulation of nutlin-3a. The intestine had an atypical profile, possibly due to biliary excretion of nutlin-3a. The blood flow-limited model did not fit well to the intestinal concentration data, limiting the ability to accurately estimate the partition coefficient for this organ.

Retinoblastoma is a tumor of the eye for which a number of orthotopic xenograft and genetic murine models have been developed [258]. Daily subconjunctival administration of nutlin-3a for 5 days was effective in a model of retinoblastoma and when combined with topotecan, an 82-fold reduction in tumor burden with no systemic or ocular side-effects was observed [24]. Our PBPK model shows that with the nutlin-3a regimen most commonly used in preclinical studies (200 mg/kg administered orally twice daily), unbound concentrations of nutlin-3a in the retina continuously exceeded the unbound IC50. However, due to poor penetration, the unbound IC50 was never achieved in the vitreous with this regimen and was achieved only transiently at 400 mg/kg twice daily, suggesting that subconjunctival dosing would be more appropriate for targeting retinoblastoma vitreous seeds [259, 260]. While the PBPK model cannot predict nutlin-3a pharmacokinetics after subconjunctival administration because the ocular absorption is not known, data from a limited experiment could be combined with the PBPK model to predict exposures in various tissues after ocular administration.

At diagnosis, 98% of neuroblastoma tumors contain wild-type p53 [209-211], and thus these patients are likely to benefit from reactivation of this pathway. Sensitivity to nutlin-3a has been shown in multiple p53 wild-type neuroblastoma cell lines [221, 228]. Treatment of a subcutaneous UKF-NB-3 rDOX20 xenograft model with twice-daily oral nutlin-3a (200 mg/kg) only partially inhibited tumor growth [43], despite good sensitivity.
(IC$_{50}$ of 5.56 μM) of these cells in vitro [228]. Because the racemic nutlin-3 mixture was used in this experiment, the assumed equivalent dose would be 100 mg/kg nutlin-3a twice daily. If unbound plasma concentrations are evaluated (since this was not an orthotopic xenograft), unbound nutlin-3a concentrations are predicted to be continually below the established IC$_{50}$ with this regimen. However, the pharmacokinetic properties of the racemic mixture are unknown, and it is possible that nutlin-3b influences the saturable elimination or plasma protein binding of nutlin-3a. For a subcutaneous or orthotopic xenograft with similar cell sensitivity, 400 mg/kg oral nutlin-3a twice daily may result in better activity, since concentrations would continuously be above the IC$_{50}$. Although the literature and our unpublished observations suggest that twice daily 200 mg/kg oral dosing is well tolerated in mice [43], further toxicity studies will need to be performed to determine whether higher dosages are tolerable.

Nutlin-3a has also demonstrated cytotoxicity in rhabdomyosarcoma cell lines [228, 249], although it has not yet been tested in a preclinical model of rhabdomyosarcoma. Our model predictions show that the standard twice-daily oral 200 mg/kg nutlin-3a regime sufficient to achieve unbound muscle nutlin-3a concentrations that were continuously above the IC$_{50}$ for the RMS-YM cell line. Nutlin-3a concentrations above the IC$_{50}$ for primary T-ALL cells [26, 41] was achieved in the plasma and spleen with a twice-daily regimen of 100 mg/kg oral nutlin-3a. However, penetration to the bone marrow was poor, and in order to target this compartment, 400 mg/kg twice-daily is recommended, which is predicted to result in unbound nutlin-3a concentrations that are above the IC$_{50}$ 77% percent of the time.

Although the model has a number of applications, it also has limitations. First, we did not perform experiments to determine the route of elimination of nutlin-3a. Preliminary unpublished observations indicate that nutlin-3a is metabolized by mouse liver microsomes, but a model with elimination only from the liver did not adequately fit the nutlin-3a plasma concentration-time data. Not modeling the elimination mechanistically could limit the ability to extrapolate the PBPK model to other species. Second, we performed all experiments in non-tumor bearing mice. Compared to normal tissues, the altered environment in tumors (e.g. vasculature, pH, interstitial fluid pressure) may lead to different local disposition. Although it was not feasible to perform pharmacokinetic studies in multiple preclinical models, data obtained from studies in tumor-bearing mice could be easily incorporated into this PBPK model. Another limitation is the assumption that the unbound fraction of nutlin-3a in tissues was equivalent to the unbound fraction in plasma. We did directly measure nutlin-3a binding in vitreous fluid, which is mostly water, but has a variety of proteins [261]. We also performed simulations at dosages beyond those used to develop the model (i.e. 400 mg/kg). Although we were able to characterize the non-linear elimination at higher plasma concentrations, it is possible that there are unknown non-linear absorption or elimination processes occurring at this higher dosage which would make these model predictions inaccurate.

In summary, we performed extensive mouse pharmacokinetic studies of nutlin-3a and developed a PBPK model, which was utilized to design nutlin-3a dosing regimens for
preclinical models of pediatric malignancies. Although there are limitations to extrapolating *in vitro* cytotoxicity data, this analysis provides a starting point for further pharmacokinetic/pharmacodynamic studies in tumor bearing mice. For models of retinoblastoma, the disposition of nutlin-3a after subconjunctival administration should be explored.
CHAPTER 5. SUMMARY AND FUTURE DIRECTIONS

The development of more effective and less toxic anti-cancer agents is a slow, high-risk, high-cost, and often high failure process. Potential new agents are emerging continuously from high throughput screening, however, only a small percentage of candidate molecules make their way from bench to bedside. Studies have shown that the success rate for the development of oncology drugs is much lower than that for drugs in other therapeutic areas [262]. In the last few years, the total capitalized cost per approved new drug molecule has risen to $0.8-1.3 billion [263-265]. For pediatric malignancies such as retinoblastoma, the small patient pool further hinders the drug development process because of the inability to conduct the clinical trials necessary to evaluate the different compounds. These difficulties necessitate better translational research, where preclinical studies can play an important role.

Since oncology drugs are usually administered in combination, it is important to evaluate the interaction of drugs. Combining agents that act synergistically instead of antagonistically to inhibit cancer cell growth can have a positive impact on patient outcome, taking into consideration that oncology drugs have narrow therapeutic windows. Here we first reported that nutlin-3a synergistically inhibited neuroblastoma cell growth in combination with topotecan and synergistically inhibited BCRP over-expressing osteosarcoma cell growth in combination with mitoxantrone. Preclinical studies from our lab [212] and Michaelis et al. [228] suggested that nutlin-3a inhibited the function of three major efflux transporters BCRP, P-gp, and MRP1. These studies suggested that further in vivo drug-drug interaction studies should be conducted because nutlin-3a may significantly increase the systemic concentrations of agents that are substrates of these transporters. This may cause severe toxicity in clinic thus extra cautious are needed. We still do not know if nutlin-3a also inhibits other efflux or uptake transporters. In addition, because of the interaction of transporters and drug metabolizing enzymes, it is also possible that nutlin-3a may inhibit drug metabolizing enzymes. In fact, our preliminary study suggested that nutlin-3a may be an inhibitor of CYP3A4 (Figure 5-1). Thus, additional preclinical studies are needed to address the interaction of nutlin-3a with CYP3A4 substrates in vitro and in vivo.

Reasons for attrition of oncology new chemical entities during clinical trials include incorrect hypothesis regarding the drug action on the disease, improper preclinical tumor model, improper clinical trial design, and drug toxicity. One additional explanation for this lack of effectiveness in translating laboratory science into efficient therapies is the scarce information on drug concentration within human tumor. The drug exposure at the target site, which is associated with efficacy, can be very different from the plasma exposure. Moreover, different tissues are likely to have different drug penetration characteristics. However, due to ethical and practical reasons, most current clinical trials do not measure drug concentrations directly in tumor or target tissues and instead rely on indirect assessment of drug exposure by measuring drug concentrations in surrogate compartments such as plasma. Thus, if one could develop an approach to reliably estimate human tumor or target tissue drug concentrations from preclinical data,
Figure 5-1. Nutlin-3a inhibits the activity of CYP3A4 *in vitro*

Study was performed using Promega p450glo CYP3A4/Luciferin-IPA Assay (Madison, WI) following manufacture instructions. High luminescence reading represents high CYP3A4 activity.
the use of drugs could be greatly improved. PK modeling and simulation is an example of such an approach. Once a preclinical model describing both plasma and target tissue/tumor concentration has been established, it can be scaled-up to humans to predict if the dosage necessary for a desirable target tissue or tumor concentration is practically achievable. If the answer is no, a "No-Go" decision should be considered to avoid trial failure and thus save precious resources. When the human plasma PK parameters are generated after either Phase 0 microdosing or after the Phase I clinical trial, the human plasma PK model can be updated to give more accurate prediction of human tissue/tumor concentration. This will allow the clinician to select more “effective” dosages, which are based upon exposures necessary for tumor effect.

In this study, we conducted developed a mouse PBPK model of nutlin-3a in the plasma and multiple tissues of therapeutic interest. This is the first study to provide comprehensive pharmacokinetic data of nutlin-3a in any species. The design of our study included both oral and intravenous dosing at several dosage levels. This method permitted the development of a robust model that accurately describes the disposition of nutlin-3a in the plasma and organs over a wide range of concentrations. The PBPK model was used to suggest dosing regimens to attain putative cytotoxic nutlin-3a exposures in target tissues for retinoblastoma, neuroblastoma, rhabdomyosarcoma, and leukemia. Once nutlin-3a plasma concentration information in human is available (either by scaling or phase 0/I clinical trial), current PBPK model can be updated to predict nutlin-3a exposure in the human tissues.

In future studies, nutlin-3a concentrations in preclinical models of pediatric tumors can be further assessed. One possible tumor to study is retinoblastoma. Nutlin-3a has shown promising efficacy in preclinical studies of retinoblastoma, and will be an excellent compound to use as a model for further PK-PD studies. Subconjunctival administration of nutlin-3a in combination with subconjunctival topotecan for 5 consecutive days reduced the tumor burden by 82-fold in rats [24]. Subconjunctival nutlin-3a in combination with systemic topotecan demonstrated significant improvement in outcome in mice bearing orthotopic retinoblastoma tumor [42]. It has been shown that the dosing route of nutlin-3a has a great impact on its ocular distribution. PK studies suggested subconjunctival administration of nutlin-3a increased the vitreous exposure by 2000-fold compared to systemic administration in non-tumor bearing mice [42]. However, the amount or extent of nutlin-3a reaching a retinoblastoma tumor in mouse models is still unknown, and it would not be feasible to obtain that data in patients. Thus, the overall goal of the proposed future study is to develop PK models of nutlin-3a disposition in retinoblastoma tumors after subconjunctival injection and systemic administration in orthotopic retinoblastoma xenograft. Both classical and PBPK model will be developed. Major challenges include determining retinoblastoma tumor blood flow, dealing with large variations of nutlin-3a concentrations in tumor, vitreous, and retina, and the structural complexity of the pharmacokinetic models. Other methodological issues to be addressed include the extremely small volume (< 8 μL per sample) of tissue samples (tumor, vitreous and retina), which could be an obstacle for both sample collection and LC-MS/MS analysis. However, when developed, these pharmacokinetic models would be the first to describe drug concentration in
retinoblastoma tumors with the ultimate goal to translate the findings to the design of clinical trials.
LIST OF REFERENCES


Fan Zhang was born in China in 1978. In 1996, Fan Zhang enrolled in Xi’an Jiaotong University, major in Clinical Medicine. In 2001, Fan received the Bachelor of Medicine degree with honor. In 2004, she received the Master of Science degree in Microbiology from the same University. She enrolled in the Ph.D. program in the Department of Pharmaceutical Sciences at the University of Tennessee Health Science Center in 2006 and received Ph.D. degree in 2011. Her dissertation research was conducted at St. Jude Children’s Research Hospital in Dr. Clinton Stewart’s lab. She is a member of the Rho Chi Honor Society.