Antiretroviral Drug Concentrations in Lymph Nodes: A Cross-Species Comparison of the Effect of Drug Transporter Expression, Viral Infection, and Sex in Humanized Mice, Nonhuman Primates, and Humans

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ABSTRACT

In a “kick and kill” strategy for human immunodeficiency virus (HIV) eradication, protective concentrations of antiretrovirals (ARVs) in the lymph node are important to prevent vulnerable cells from further HIV infection. However, the factors responsible for drug distribution and concentration into these tissues are largely unknown. Although humanized mice and nonhuman primates (NHPs) are crucial to HIV research, ARV tissue pharmacology has not been well characterized across species. This study investigated the influence of drug transporter expression, viral infection, and sex on ARV penetration within lymph nodes of animal models and humans. Six ARVs were dosed for 10 days in humanized mice and NHPs. Plasma and lymph nodes were collected at necropsy, 24 hours after the last dose. Human lymph node tissue and plasma from deceased patients were collected from tissue banks. ARV, active metabolite, and endogenous nucleotide concentrations were measured by liquid chromatography–tandem mass spectrometry, and drug transporter expression was measured using quantitative polymerase chain reaction and quantitative targeted absolute proteomics. In NHPs and humans, lymph node ARV concentrations were greater than or equal to plasma, and tenofovir diphosphate/deoxyadenosine triphosphate concentration ratios achieved efficacy targets in lymph nodes from all three species. There was no effect of infection or sex on ARV concentrations. Low drug transporter expression existed in lymph nodes from all species, and no predictive relationships were found between transporter gene/protein expression and ARV penetration. Overall, common preclinical models of HIV infection were well suited to predict human ARV exposure in lymph nodes, and low transporter expression suggests primarily passive drug distribution in these tissues.

SIGNIFICANCE STATEMENT

During human immunodeficiency virus (HIV) eradication strategies, protective concentrations of antiretrovirals (ARVs) in the lymph node prevent vulnerable cells from further HIV infection. However, ARV tissue pharmacology has not been well characterized across preclinical species used for HIV eradication research, and the influence of drug transporters, HIV infection, and sex on ARV distribution and concentration into the lymph node is largely unknown. Here we show that two animal models of HIV infection (humanized mice and nonhuman primates) were well suited to predict human ARV exposure in lymph nodes. Additionally, we found that drug transporter expression was minimal and—along with viral infection and sex—did not affect ARV penetration into lymph nodes from any species.

Introduction

Advances in antiretroviral therapy (ART) potency and dosing have transformed human immunodeficiency virus (HIV) from a fatal diagnosis to a chronic condition, with patient life expectancy now comparable to that of an uninfected population (Samji et al., 2013). Despite this, challenges in treating and curing HIV remain (Lewin and Rouzioux, 2011). Although the majority of patients adherent to ART show undetectable viral replication in blood plasma, viral rebound occurs upon interruption of therapy (Chun et al., 1999). One theoretical cause of viral rebound is low-level viral replication within tissues (Natarajan et al., 1999; Buzón et al., 2011), caused by inadequate antiretroviral (ARV) penetration (Horiike et al., 2012; Fletcher et al., 2014, 2018). Low ARV...
penetration may also present challenges in protecting cells from infection during “kick and kill” strategies for HIV eradication (Lewin and Rouzioux, 2011).

In the area of HIV prevention, investigations into drug concentrations in colorectal and female genital tract tissues have shown highly variable ARV penetration (Dumond et al., 2007, 2009; Kwara et al., 2008; Else et al., 2011; Patterson et al., 2011; Thompson et al., 2015; Cottrell et al., 2016). However, considerably less drug exposure data are available for the putative lymph node reservoir, despite the fact that these tissues are central to the pathology of HIV (North et al., 2010; Dimopoulos et al., 2017). In the small number of published studies, results are conflicting. ARV lymph node concentrations range from 2-fold higher to 1000-fold lower than those in plasma (Kinman et al., 2003; Solas et al., 2003; Fletcher et al., 2014). Furthermore, the methods used to investigate drug penetration in lymph nodes are also varied; some studies have used tissue homogenate (Solas et al., 2003), whereas others have isolated lymph node mononuclear cells (LNMCs) (Kinman et al., 2003; Fletcher et al., 2014).

Because human lymph node collection is invasive, preclinical models such as HIV-infected humanized mice and nonhuman primates (NHPs) with reverse transcriptase–sminian/HIV may be informative for characterizing the pharmacokinetics of ARVs in lymph nodes. Preclinical models have been used to assess viral dynamics in tissues and curative ART strategies (Denton and Garcia, 2011; Evans and Silvestri, 2013), but a cross-species comparison of ARV concentrations in putative tissue reservoirs is still needed. Furthermore, to understand concentration-response relationships of ARVs in tissues, it is important to investigate relevant biologic and disease-related factors that may influence these concentrations, such as drug transporters, viral infection, and sex (Antonelli et al., 1992; Speck et al., 2002; Fletcher et al., 2004; Giraud et al., 2010; Alam et al., 2016; Kis et al., 2016).

In this study, we aimed to quantify ARV concentrations and these aforementioned influencing factors using lymph nodes from humanized mice, NHPs, and deceased HIV-positive patients. We believe that these data improve our understanding of ARV pharmacology in the lymph node—an important but pharmacologically less characterized HIV tissue reservoir.

Materials and Methods

Animal Studies and Lymph Node Collection. A detailed description of the animal studies was published previously (Thompson et al., 2017) and is summarized here. This study used three common HIV preclinical models from two species: hu-HSC-Rag (RagHu) humanized mice (n = 36), bone marrow–liver-thymus (BLT) humanized mice (n = 13), and rhesus macaques (NHPs) (n = 18). All humanized mice and six NHPs were female. Aggressive interactions are common between male mice within the same cage, but isolating mice in separate cages can be inefficient and costly. For these reasons, we chose to use only female mice that could be safely kept in groups to minimize cage needs while maximizing animal well-being (Van Loo et al., 2003). RagHu mice (n = 18), BLT mice (n = 7), and NHPs (n = 10) were infected for 6 weeks with HIVRAL, D7, HIVmac239, and reverse transcriptase–sminian/HIV, respectively (North et al., 2010). The remaining animals were uninfected (RagHu, n = 18; BLT, n = 6; and NHP, n = 8). After the infection period, animals were dosed for 10 days to steady state and necropsy was performed 1 day after the last dose. Doses were chosen based on previously reported effective treatment regimens in these animal models (Denton et al., 2010; Neff et al., 2010; Shytaj et al., 2012; Massaud et al., 2013; Veselinovic et al., 2014) and are reported in the Supplemental Methods. RagHu mice were dosed with efavirenz (EFV) only (n = 12), atazanavir (ATZ) only (n = 12), or a combination of tenofovir (TFV), emtricitabine (FTC), raltegravir (RAL), and maraviroc (MVC) (n = 12). All BLT mice received the five drug combination of TFV/FTC/RAL/MVC/ATZ (n = 13). Finally, NHPs were dosed with TFV/FTC/EFV/RAL (n = 9) or TFV/FTC/MVC/ATZ (n = 9). Further details on animal dosing, infection, and tissue collection can be found in the Supplemental Methods, Supplemental Fig. 1, and Supplemental Table 1.

At necropsy, plasma and lymph nodes were collected from all animals and snap frozen. For both RagHu and BLT mice, lymph node type was not determined; for NHPs, four lymph nodes (mesenteric, axillary, inguinal, and iliac) per animal were collected. Mouse and NHP tissue was divided for use in ARV and transporter expression analyses, with matched data obtained when possible (Supplemental Table 2). All animal studies were performed in accordance with Institutional Animal Care and Use Committee protocols from the University of North Carolina at Chapel Hill (protocol 15-168), Colorado State University (protocol 16-6998A), and the University of California, Davis (protocol 18345).

Human Lymph Node Collection. Human plasma and lymph node samples were obtained by the National Research Disease Interchange and the National NeuroAIDS Tissue Consortium from deceased HIV-positive patients who consented to organ donation prior to death or whose families consented to organ donation immediately after the patients’ death. To screen for subjects who were adherent to ART or received ARV dosing premortem (and therefore most likely to have quantifiable tissue ARV concentrations), plasma samples were obtained based on desired subject characteristics such as tissue availability, viral load, and ART regimens containing TFV, FTC, EFV, RAL, MVC, or ATZ. Plasma samples were analyzed in our laboratory by liquid chromatography–tandem mass spectrometry (LC-MS/MS), and only subjects with measurable ARV plasma concentrations were included in the tissue request. Ultimately, lymph node sections from 13 subjects (two female) were requested, and each subject’s ART regimen contained at least one drug of interest (eight TFV, four FTC, seven EFV, three RAL, two ATZ, and zero MVC). Human lymph node tissue was divided for use in ARV and transporter expression analyses, with matched data obtained when possible (Supplemental Table 2). Details on subject demographics, virology, ART dosing/regimens, and preliminary plasma ARV concentrations can be found in Supplemental Table 3.

ARV Concentration and Tissue Penetration Analyses. Concentrations of all six ARVs (TFV, FTC, EFV, RAL, MVC, and ATZ) were analyzed in plasma and lymph node tissue using LC-MS/MS. Additionally, the intracellular active metabolites tenofovir diphosphate (TFVdp) and emtricitabine triphosphate (FTCtp), and their respective endogenous nucleotides, deoxyadenosine triphosphate (dATP) and deoxycytidine triphosphate (dCTP), were analyzed in lymph node tissue.

ABBREVIATIONS: ART, antiretroviral therapy; ARV, antiretroviral; ATZ, atazanavir; BLQ, below the limit of quantification; BLT, bone marrow–liver-thymus humanized mice; dATP, deoxyadenosine triphosphate; dCTP, deoxycytidine triphosphate; EC50, 90% effective concentration; EFV, efavirenz; ENT1, equilibrative nucleoside transporter 1; FTC, emtricitabine; FTCtp, emtricitabine triphosphate; GAPDH, glyceraldehyde 3-phosphate dehydrogenase; HIV, human immunodeficiency virus; LC-MS/MS, liquid chromatography–tandem mass spectrometry; LLOQ, lower limit of detection; LNMC, lymph node mononuclear cell; MRP, multidrug resistance–associated protein; MSI, mass spectrometry imaging; MVC, maraviroc; NHP, nonhuman primate; P-gp, P-glycoprotein; RagHu, hu-HSC-Rag; RAL, raltegravir; TFV, tenofovir; TFVdp, tenofovir diphosphate; TPR, tissue penetration ratio.
Plasma and tissue samples were extracted by protein precipitation using stable, isotope-labeled internal standards. Extracts were analyzed by a Shimadzu high-performance liquid chromatography system with an API 5000 mass spectrometer (SCIEX, Framingham, MA) detector equipped with a TurbolonSpray interface. The lower limit of quantitation was 1 ng/ml for plasma and 0.002 ng/ml (FTC, MVC), 0.005 ng/ml (EFV, RAL, ATZ), 0.01 ng/ml (TFV), 1.11 ng/ml (dATP, dCTP), and 0.22 ng/ml (TFVdp, FTCtp) for tissue. Assay precision and accuracy was within 15%.

At least 30 mg lymph node tissue was homogenized using a Preceelys Tissue Homogenizer (Bertin Technologies, Montigny-le-Bretonneux, France) in 1 ml of 70:30 acetonitrile:1 mM ammonium phosphate (pH 7.4) and extracted by protein precipitation as before. TFV and FTC were analyzed on a Waters Atlantis T3 (50 mm × 2.1 mm, 3 μm particle size) column. EFV, RAL, MVC, and ATZ were separated using an Agilent Pursuit XRs 3 Diphenyl (50 mm × 2 mm, 5 μm particle size) high-performance liquid chromatography column. TFVdp, FTCtp, dATP, and dCTP were analyzed on a Thermo BioBasic AX column. Concentrations were converted from nanograms per millilitre to nanograms per gram based on a tissue density of 1.06 g/ml.

Tissue penetration ratios (TPRs) at the end of the dosing interval were calculated for each ARV by dividing lymph node concentrations by plasma concentrations. We also calculated active metabolite to endogenous nucleotide ratios, since the efficacy of the active metabolite depends on its concentration relative to the endogenous nucleotide it replaces during reverse transcription (Anderson et al., 2011).

**Protein Expression Analysis.** Protein concentrations of five efflux and three uptake transporters were measured by quantitative targeted absolute proteomics (Fallon et al., 2013). Transporters were chosen based on their relevance to ARV disposition, putative expression in lymph nodes, or previous investigation in other reservoir tissues (Supplemental Table 4) (Minuesa et al., 2011; Zhou et al., 2013; Nicol et al., 2014; Huang et al., 2016; Thompson et al., 2017). Approximately 100 mg lymph node tissue was homogenized in 1.3 ml hypotonic buffer [10 mM NaCl, 1.5 mM MgCl2, 10 mM Tris HCl, pH 7.4, and 150 μM Complete Protease Inhibitor Solution (Sigma-Aldrich, St. Louis, MO)] using a Preceelys Tissue Homogenizer, and 10–30 μg membrane protein was isolated as previously described (Fallon et al., 2016; Thompson et al., 2017). Membrane protein was dried down and reconstituted with 50 mM ammonium bicarbonate buffer plus 40 mM dithiothreitol, 10% sodium deoxycholate, and 10 μl β-casein (0.1 μg/μl). Samples were then reduced for 40 minutes at 60°C followed by addition of 135 mM iodoacetamide and incubation in the dark for 30 minutes at room temperature. One percent of stable isotope labeled peptide standards (Theracode JPT Inc., Acton, MA) was added to the samples, followed by digestion with 25 μl trypsin (0.1 μg/μl) at 37°C (Promega, Madison, WI). Digestion was interrupted after 18 hours with 10% trifluoroacetic acid, and samples were extracted using solid phase extraction on 33-m particle size) columns. Samples were then reduced for 40 minutes at 60°C followed by addition of 135 mM iodoacetamide and incubation in the dark for 30 minutes at room temperature. One percent of stable isotope labeled peptide standards (Theracode JPT Inc., Acton, MA) was added to the samples, followed by digestion with 25 μl trypsin (0.1 μg/μl) at 37°C (Promega, Madison, WI). Digestion was interrupted after 18 hours with 10% trifluoroacetic acid, and samples were extracted using solid phase extraction on 33-μm polymeric reversed phase extraction columns (Phenomenex, Torrance, CA). After final dry-down and reconstitution in 98% formic acid (0.1%) plus 2% acetonitrile, around 0.06–0.12 μg microsomal protein was loaded onto a C18 trap column connected to a BEH130 C18 (150 μm × 100 mm, 1.7 μm particle size) main separation column. Sample analysis was performed on a nanoACQUITY system (Waters, Milford, MA) coupled to a Qtrap 5500 mass spectrometer (SCIEX) equipped with a Nanospray III source. Analyst 1.5 and MultiQuant 2.0 software (SCIEX) were used for multiple reaction monitoring data acquisition and analysis. Peak area ratios of unlabeled/stable isotope labeled peptides were determined using the sum of two multiple reaction monitoring values. The lower limit of detection (LLOD) for the peptides was 0.1 pmol/mg protein, and 50 μg NHP liver homogenate was used as a positive control.

**Gene Expression Analysis.** Gene expression of the same eight transporters was measured by quantitative polymerase chain reaction. Approximately 30 mg tissue was homogenized using a Preceelys Tissue Homogenizer and RNA was extracted using a Qiagen RNeasy Fibrous Tissue Mini Kit (Qiagen, Valencia, CA). The VILO Superscript cDNA Synthesis Kit (Thermo Fisher, Waltham, MA) was used to reverse transcribe 200 ng RNA. A 10-cycle preamplification was performed, followed by 40 cycles of quantitative polymerase chain reaction using TaqMan primers and probes (Supplementary Table 5) on a QuantStudio 6 system (Life Technologies, Waltham, MA). Samples were run in triplicate, and transporter expression was normalized to the housekeeping gene glyceraldehyde 3-phosphate dehydrogenase (GAPDH) using the 2^(-ΔΔCt) method (Schmittgen and Livak, 2008). The LLOD was a 0.0001-fold change over GAPDH.

**Statistical Analysis.** In all species, ARV concentrations, gene expression values, or transporter protein concentrations below the limit of detection or below the limit of quantitation (BLQ) of the assay were imputed at half the lower limit of quantitation or LLOD, respectively. Relationships between plasma and lymph node drug concentrations were assessed using linear regression. Comparisons between species, infection status, and sex were performed using the Kruskal–Wallis one-way ANOVA with Dunn’s method for multiple comparisons, and relationships between transporter expression and TPRs were investigated using multiple linear regression. Data were analyzed using SigmaPlot 13.0 (Systat Software Inc., San Jose, CA) with a significance level of P < 0.05.

**Results**

There were no statistically significant differences in ARV concentrations between the four lymph nodes collected from each NHP (either uninfected or infected) (Supplemental Fig. 2), so data from all lymph nodes were averaged to generate one median value for each uninfected or infected animal. Plasma and lymph node ARV concentrations were not significantly different between BLT and RagHu mice, so data from both humanized mouse species were pooled for the remaining analyses (Supplemental Fig. 3). One female macaque dosed with TFV/FTC/EFV/RAL developed liver failure, resulting in drug concentrations 17- to 260-fold higher than other animals in the dosing group. This NHP was excluded from ARV concentration/penetration analyses but was included in drug transporter comparisons; transporter expression was not significantly different from other animals in the dosing group. Additionally, to avoid overestimating lymph node ARV penetration, mice and NHPs with BLQ plasma concentrations but measurable lymph node concentrations were excluded from analyses. These sample size adjustments are reflected in Supplemental Table 2.

**ARV Concentrations in Plasma and Lymph Nodes.**

Several species-dependent trends in ARV concentrations and lymph node penetration relative to plasma were noted. Median ARV plasma concentrations were generally lowest in mice (14% BLQ) and highest in humans, with a few exceptions: FTC and TFV were lowest in NHPs rather than mice, and RAL was highest in NHPs rather than humans; however, only one human plasma sample was available for RAL analysis (Fig. 1A). In lymph nodes, median ARV concentrations followed the same species trends seen in plasma, with 46% of mouse concentrations BLQ. The only notable differences were in TFV (highest in NHP's rather than humans) and RAL (highest in humans) (Fig. 1B). TPRs were calculated using matched lymph node and plasma concentrations. Median NHP lymph node TPRs ranged from 3-fold lower (RAL) to 25-fold higher (MVC) than plasma (Fig. 2). Compared with other species, NHP TPRs were highest for all drugs except FTC and RAL. Median TPRs ranged from 0.0002 (RAL) to 8 (FTC) for mice and from 0.8 (ATZ) to 2 (EFV) for humans.
Excluding BLQ concentrations in mice, there were several significant relationships between lymph node and plasma concentrations: FTC ($r^2 = 0.73, P < 0.001$), TFV ($r^2 = 0.03$), EFV ($r^2 = 0.73, P < 0.001$), RAL ($r^2 = 0.004, P < 0.05$), MVC ($r^2 = 0.40, P < 0.05$), and ATZ ($r^2 = 0.55, P < 0.01$).

**Active Metabolite and Endogenous Nucleotide Concentrations and Ratios.** Active metabolite concentrations showed greater between-species variability than parent drugs, particularly for TFVdp. Similar to the pattern of TFV TPRs, median TFVdp concentrations were highest in NHPs (Fig. 3A). On a molar basis, lymph nodes from all species contained more parent drugs than active metabolites, but NHPs had the greatest proportion of FTC converted to FTCtp (4.8%; mouse = 0.2%, human = 1.1%) and TFV converted to TFVdp (6.4%; mouse = 0.1%, human = 0.2%) (Table 1). Concentrations of the endogenous nucleotides dCTP and dATP were highest in NHP lymph nodes but were not significantly different across species (Fig. 3A). When comparing endogenous nucleotides to corresponding active metabolites, only NHP dATP concentrations were significantly different from TFVdp concentrations ($P < 0.05$). Therefore, after normalizing active metabolite concentrations to endogenous nucleotides, TFVdp/dATP ratios were significantly higher in NHPs, but FTCtp/dCTP ratios were similar across species (Fig. 3B).

**Effect of Viral Infection and Sex on ARV Penetration in Lymph Nodes.** No differences in lymph node drug penetration due to viral infection (Fig. 4) or sex (Fig. 5) were found in any species. There were also no trends in the variability of ARV penetration between uninfected and infected animals or males and females.

**Drug Transporter Expression in Lymph Nodes.** Lymph node drug transporter expression was low across all species. Gene expression of all eight transporters was consistently lower than the housekeeping gene GAPDH in both mice and NHPs (Fig. 6A). Median equilibrative nucleoside transporter 1 (ENT1) gene expression was highest overall for both species. Mouse gene expression was higher than NHP for all transporters but multidrug resistance–associated protein 2 (MRP2). Transporter protein expression was also low in NHP and human lymph nodes; breast cancer resistance protein, ENT1, and organic cation transporter 3 were the only NHP transporters with median concentrations above the limit of detection (94%, 100%, and 94% of samples above LLOD, respectively), and median concentrations of all eight transporters were below the limit of detection in humans (Fig. 6B).

**Effect of Drug Transporter Expression on ARV Lymph Node Penetration.** Few predictive relationships were found between ARV TPRs and drug transporter expression. Higher mouse and NHP gene expression of MRP2 was predictive of greater TFV penetration in lymph nodes ($P = 0.002, r^2 = 0.27$) (Supplementary Table 6), and greater ATZ penetration in NHPs and humans was predicted by an increase in combined MRP1 and P-glycoprotein (P-gp) protein expression ($P < 0.001, r^2 = 0.99$) (Supplementary Table 7).
Discussion

This study is the first to evaluate ARV exposure in lymph nodes of three species. In this investigation, we determined that there was no effect of acute HIV infection or sex on ARV lymph node concentrations. However, we did find several interesting species-dependent differences in drug exposure. In mice, EFV, RAL, MVC, and ATZ penetration was low and variable: between 33% (EFV) and 83% (RAL) of lymph node concentrations were BLQ. No relationship was found between the amount of mouse lymph node analyzed and tissue concentrations, indicating that BLQ concentrations are not the result of small tissue quantities and LC-MS/MS sensitivity. There were also no animal-specific trends in tissue concentrations: previous analyses found no BLQ concentrations in brain tissue from any mouse, whereas 96% of ileum and 87% of rectum concentrations were BLQ in the same animals (Thompson, 2017; Srinivas et al., 2018). When considering only mice with quantifiable lymph node concentrations, drug penetration was generally similar between mice, NHPs, and humans.

Similar to the pattern of TFV penetration ratios, TFVdp concentrations and TFVdp/dATP ratios were significantly higher in NHPs. Previous work in NHPs and humans has shown that in target lymphoid organs, the concentration ratio of TFVdp to the endogenous substrate dATP may be a better predictor of efficacy than TFVdp concentrations alone (García-Lerma et al., 2011; Cottrell et al., 2016). Using an in vitro enzymatic inhibition assay, investigators have shown that a TFVdp/dATP ratio ≥1 corresponds to 100% reverse transcriptase inhibition in NHPs (García-Lerma et al., 2011). In our study, median NHP TFVdp/dATP ratios were 34, well above an efficacy target of 1. Similarly, a predictive pharmacokinetic/pharmacodynamic model in humans found a 90% effective TFVdp/dATP concentration ratio (EC$_{90}$) of 0.29 for protecting CD4+ T cells from viral challenge (Cottrell et al., 2016); in our study, this efficacy target was exceeded by NHPs and humans and almost achieved in mice (median ratios of 34, 1, and 0.2, respectively). These data show that despite a species difference in TFVdp concentrations, both preclinical species meet previously investigated human TFVdp/dATP efficacy targets in lymph node tissue.

A previous study reported that TFVdp concentrations in human LNMCs were significantly lower than in peripheral blood mononuclear cells and that these low concentrations were associated with HIV replication (Fletcher et al., 2014). Indeed, it is well known that even during suppressive ART, viral RNA and DNA can persist in lymph nodes and other tissues from NHPs and humans (North et al., 2010; Deleage et al., 2016; Lamers et al., 2016; Estes et al., 2017), and there may be evidence of ongoing HIV replication within the lymph nodes of ART-suppressed patients (Lorenzo-Redondo et al., 2016; Halvas et al., 2019). However, in contrast to previous LNMC studies—which do not account for drug lost during cell isolation and may underestimate concentrations—we found that lymph node TFV concentrations are higher than or equal to plasma, and TFVdp/dATP ratios meet or exceed the EC$_{90}$ target to inhibit HIV reverse transcription.

Yet the concentrations we have presented are an averaged concentration throughout the lymph node. It has been previously demonstrated that drug distribution within a tissue can be variable (Fischman et al., 1998), and we have shown heterogeneous EFV concentrations across five different putative tissue reservoirs using mass spectrometry imaging (MSI) (Thompson et al., 2015). Recently, we also saw little colocalization of viral RNA with efficacious ARV concentrations in the brain (3% overlap) (Srinivas et al., 2018). Therefore, an ARV’s location within tissue is as important as overall concentration. We are continuing to evaluate lymph node distribution by quantitative MSI to address this issue.

In addition to ARV concentrations, this study was also the first to examine cross-species expression of relevant ARV transporters in lymph nodes. Overall, transporter expression in lymph nodes is comparable to what we have seen in brain tissue (Srinivas, 2018) but lower than other HIV reservoirs such as the testis, ileum, and rectum (Huang et al., 2016; Thompson et al., 2017). The most highly expressed

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TABLE 1

<table>
<thead>
<tr>
<th>ARV</th>
<th>Mouse</th>
<th>NHP</th>
<th>Human</th>
</tr>
</thead>
<tbody>
<tr>
<td>FTC</td>
<td>99.8 (77.4, 100)</td>
<td>95.2 (79, 100)</td>
<td>98.9 (21, 100)</td>
</tr>
<tr>
<td>FTCtp</td>
<td>0.2 (0, 22.6)</td>
<td>4.8 (0, 21)</td>
<td>1.1 (0, 97.9)</td>
</tr>
<tr>
<td>TFV</td>
<td>99.9 (95.6, 100)</td>
<td>93.6 (64.2, 100)</td>
<td>99.8 (89.3, 100)</td>
</tr>
<tr>
<td>TFVdp</td>
<td>0.1 (0, 4.4)</td>
<td>6.4 (0, 35.8)</td>
<td>0.2 (0, 10.7)</td>
</tr>
</tbody>
</table>
transporter by gene and protein methods was ENT1. This aligns with previous RNA expression and immunocytochemistry data for ENT1 in lymphoblastic T-cell lines and CD4+ T cells isolated from HIV-negative donors (Minuesa et al., 2008). ENT1 is responsible for the cellular uptake of nucleosides such as adenosine and uridine, which are important for immunostimulatory responses and lymphocyte proliferation (Goodman and Weigle, 1983, 1984); therefore, increased ENT1 expression in lymph nodes is not unexpected.

To determine whether even low-level transporter expression could influence drug distribution in the lymph node, we used multilinear regression to assess relationships between ARV penetration ratios and transporter gene/protein expression across species, including any potential link between ENT1 expression and TFV (an adenosine analog) (Viread, 2001). Although the positive associations between TFV versus MRP2 (in mice and NHPs) and ATZ versus MRP1 + P-gp (in NHPs and humans) are significant, these relationships were primarily driven by the number of below the limit of detection transporter concentrations and are inconsistent with the intracellular-to-extracellular direction of MRP1, MRP2, and P-gp efflux transporters. Taken together, generally low drug transporter expression in the lymph node, and lack of physiologically relevant predictive relationships between transporters and ARV penetration, suggests that drug distribution in the lymph node may occur by passive, diffusion-based mechanisms—contributing to the heterogeneous ARV distribution we have seen in the lymph node for EFV (Thompson et al., 2015).

There are several limitations in our assessment of ARV tissue penetration. One important consideration is our use of steady-state plasma and tissue concentrations obtained at the end of the dosing interval to estimate TPRs. This approach may overestimate drug penetration into lymph nodes, since the difference in drug concentrations in these two compartments is likely greatest at the end of the dosing interval. A better method of estimating tissue penetration is by dividing lymph node and plasma areas under the concentration time curves (AUC_tissue/AUC_plasma), which can be determined by repeated tissue sampling. However, in our animal studies, we opted to start collecting a set of whole lymph nodes at the end of the dosing interval. A second possible limitation is blood contamination within lymph node homogenate, although no blood was observed during visual inspection of the tissue.

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**Fig. 4.** Differences in ARV TPRs in uninfected (gray boxes) and infected (red boxes) mouse (A) and NHP (B) lymph nodes. Boxes are the 25th percentile, median, and 75th percentile. Whiskers are the fifth and 95th percentiles, and dots are outliers. The dashed line indicates equal drug concentrations in lymph node and plasma. Sample sizes for each drug in each species are given in Supplemental Table 2. Infection differences were analyzed using Kruskal–Wallis one-way ANOVA with Dunn’s method correction for multiple comparisons.

**Fig. 5.** Differences in ARV TPRs in female (pink boxes) and male (blue boxes) NHP (A) and human (B) lymph nodes. Boxes are the 25th percentile, median, and 75th percentile. Whiskers are the fifth and 95th percentiles, and dots are outliers. Dashed lines indicate equal drug concentrations in lymph node and plasma. Sample sizes for each drug in each species are given in Supplemental Table 2. Sex differences were analyzed using Kruskal–Wallis one-way ANOVA with Dunn’s method correction for multiple comparisons.
Using MSI we have analyzed lymph node sections for heme (a blood marker) and ARVs and found little colocalization of the two (≤25% overlap) (unpublished data). We will continue these analyses in the future to correct for blood contamination in the lymph node. Finally, lymph node ARV concentrations reported here include both protein bound and unbound fractions. Preliminary studies in a small set of NHP samples suggest that RAL, MVC, and ATZ binding is minimal in lymph nodes (<50% bound), whereas EPV binding is comparable to plasma (~97% bound) (unpublished data). Further binding analyses are ongoing.

Additionally, there were several limitations to our drug transporter analyses. We were unable to compare all tissues for gene and protein expression. Although protein expression is a more relevant measure of transporter presence in tissue (Ohtsuki et al., 2012), the extremely small size (15–30 mg) of the mouse lymph nodes was prohibitive for transporter protein analyses, which requires at least 100 mg tissue before initial homogenization. Therefore, we opted to evaluate gene expression in mice and NHPs. Since transporter expression was low across all species by both methods, it is unlikely that any appreciable transporter protein concentrations would have been found in mice even with an adequate supply of tissue. Also, our gene and protein analyses do not account for transporter activity. Future immunohistochemistry analysis of transporter localization in lymph nodes will investigate whether the majority of transporters are localized to plasma membranes, nuclear membranes, or within cytosol (where they would be functionally inactive) (Giacomini et al., 2010).

In conclusion, we measured ARV penetration and drug transporter expression in lymphated mouse, NHP, and human lymph nodes to further characterize species differences in ARV pharmacology within this putative HIV reservoir. We also investigated the effect of infection status and sex on ARV penetration, and we characterized the relationship between drug transporter expression and ARV penetration in the lymph node. Most notably, in NHPs and humans we found that lymph node ARV exposure was greater than or equal to plasma, and lymph node concentrations (with the exception of TFV and RAL) were well predicted by those in plasma. We also found that mouse and NHP TFVdp/dATP ratios met or exceeded human EC90 efficacy targets in lymph nodes. There was no effect of infection or sex on lymph node ARV penetration in any species. We also found low drug transporter expression in lymph nodes from all species and no predictive relationships between transporter gene or protein expression and ARV exposure, indicating that drug distribution in lymph nodes may occur through passive mechanisms. Our future research will focus on other physiologic (protein binding, tissue fibrosis) or physiochemical (molecular weight, lipophilicity) factors that may contribute to altered drug penetration and distribution in this reservoir. Along with the findings presented here, these data will have important implications for future preclinical and clinical drug development studies of HIV therapies targeted to lymph nodes.

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Authorship Contributions

Participated in research design: Fallon, Luciw, Garcia, Akkina, Smith, Kashuba.


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References


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