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In Vivo and in Vitro Proteome Analysis of Human Immunodeficiency Virus (HIV)-1-infected, Human CD4+ T Cells

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Host-directed therapies against HIV-1 are thought to be critical for long term containment of the HIV-1 pandemic but remain elusive. Because HIV-1 infects and manipulates important effectors of both the innate and adaptive immune system, identifying modulations of the host cell systems in humans during HIV-1 infection may be crucial for the development of immune based therapies. Here, we quantified the changes of the proteome in human CD4+ T cells upon HIV-1 infection, both in vitro and in vivo. A SWATH-MS approach was used to measure the proteome of human primary CD4+ T cells infected with HIV-1 in vitro as well as CD4+ T cells from HIV-1-infected patients with paired samples on and off antiretroviral treatment. In the in vitro experiment, the proteome of CD4+ T cells was quantified over a time course following HIV-1 infection. 1,725 host cell proteins and 4 HIV-1 proteins were quantified, with 145 proteins changing significantly during the time course. Changes in the proteome peaked 24 h after infection, concomitantly with significant HIV-1 protein production. In the in vivo branch of the study, CD4+ T cells from viremic patients and those with no detectable viral load after treatment were sorted, and the proteomes were quantified. We consistently detected 895 proteins, 172 of which were considered to be significantly different between the viremic patients and patients undergoing successful treatment. The proteome of the in vitro-infected CD4+ T cells was modulated on multiple functional levels, including TLR-4 signaling and the type 1 interferon signaling pathway. Perturbations in the type 1 interferon signaling pathway were recapitulated in CD4+ T cells from patients. The study shows that proteome maps generated by SWATH-MS indicate a range of functionally significant changes in the proteome of HIV-infected human CD4+ T cells. Exploring these perturbations in more detail may help identify new targets for immune based interventions. Molecular & Cellular Proteomics 16: 10.1074/mcp.M116.065235, S108–S123, 2017.

Therapeutic approaches against HIV-1 are mired in dichotomy. On the one hand, drugs targeting the virus are turning HIV infection successfully into a therapeutically manageable chronic disease, blocking transmission and increasing survival dramatically (1). On the other hand, host-targeted approaches such as vaccines or attempts to repair or boost the immune system failed without exception (2, 3). However, it is generally assumed that vaccines and sterilizing treatment are prerequisites to eradicate HIV-1, because lifelong treatment is difficult to implement, costly, and has side effects (4).

Current antiretroviral therapy stops viral replication but cannot kill latently infected cells. Therefore, other approaches are needed that can reverse HIV-1 latency or increase innate or adaptive immune responses to kill latently infected cells. Because the virus requires host cells for replication and survival, it needs to interact and modify the host environment to its advantage. Although these complex interactions have been extensively studied on a transcriptional level (5, 6), the knowledge about the changes on the protein level are sparse (7, 8). Therefore, more knowledge on the effects of HIV-1 on host proteome networks in human cells infected by the virus is needed to better understand virus-host interactions. It has been estimated that genetic determinants of the virus are responsible for 30–50% of the variability of the viral set point, a surrogate marker for disease progression and compared with 20% of the host genome (9). Thus, exploring host networks in clinically well characterized human individuals infected with HIV-1 is crucial for the advancement of host-targeted therapies (10). On a transcriptome level, the networks of HIV-1-infected CD4+ T cells have already been studied both in vitro (5) and in vivo (11, 12). However, the changes detectable on the transcriptome level are largely...
driven by viral replication. Therefore, they are not ideal for the discovery of mechanisms of viral control (11). In contrast, proteins are the main molecular effectors of the cell and are at the functional interface between virus and host. Analysis of the proteome may therefore be useful to detect new mechanisms associated with control of the virus.

Mass spectrometry (MS) has increasingly become the method of choice for analysis of complex protein samples, both qualitatively and quantitatively (13). We have recently developed SWATH-MS, a technique that combines the high quantitative accuracy of targeted proteomics with the broader coverage achievable with discovery proteomics. In essence, SWATH-MS is a massively parallel targeted mass spectrometric strategy that requires the a priori generation of spectral libraries that are then used to identify and quantify query peptide in the acquired datasets (14, 15). SWATH-MS provides selected reaction monitoring-like performance in terms of reproducibility, quantitative accuracy, data completeness, and dynamic range (16). Furthermore, and unlike selected reaction monitoring, SWATH-MS can quantify an unlimited number of target peptides as long as they have been previously observed by DDA1 (15).

MS approaches have been used previously to quantify the changes in the proteome of T cell lines and macrophages upon infection with HIV-1 (7, 8). However, the proteome of the main target cell of HIV-1, the human CD4+ T cell, has not been assessed yet.

In this study, we describe the results of the exploratory study in which the proteome of human CD4+ T cells, the most important target cell for HIV-1, is quantified to detect the changes associated with HIV-1 infection. By infecting human CD4+ T cells in vitro and following the effects of the infection on the host proteome over time and by assessing the proteome differences in paired samples from viremic and subsequently treated patients with no detectable viral load, we aimed to cover the changes of the CD4+ T cell proteome associated with HIV-1 infection in both in vitro and in human individuals. The data re-iterate the central role for type 1 interferon during HIV-1 infection and suggest a possibly novel role for TLR-4 signaling. Finally, the changes in the proteome during in vitro and the in vivo HIV infection are to large extent dissimilar, except for significant enrichment of type 1 interferon signaling upon functional enrichment analysis.

PATIENTS AND METHODS

Patients—10 HIV-1-infected individuals were enrolled from the longitudinal Zurich Primary HIV-1 Infection Study (ZPHI), which is an open label, non-randomized, observational, single-center study (www.clinicaltrials.gov, ID NCT00537966) (17). Blood samples at two different time points of each patient were investigated. At time point 1, the patients were not treated and had HIV-1 detectable. At time point 2, the patients were treated and had no detectable viral load for a minimum of 6 months. For patient details, see Table I.

Ethics Statement—The ethic committee of the University Hospital Zurich approved the study protocol and a written informed consent was obtained from all patients. Buffy coats from healthy donors were received from the Blood Donation Service Zurich, Swiss Red Cross, Switzerland. Written in-
formed consent was obtained for the use of buffy coats for science, which were not needed for medical purposes by the Blood Donation Center.

CD4+ T Cell Culture and Infection—Peripheral blood mononuclear cells (PBMCs) were isolated from buffy coats, as described previously (18), and enriched for CD4+ T cells by magnetic microbeads according to the manufacturer’s protocol (Miltenyi Biotec). After a three-way stimulation (phytohemagglutinin 5.0 μg/ml, phytohemagglutinin 0.5 μg/ml, and Oct3) for 3 days, the cells were infected with HIV-1JR-FL by spinoculation for 2 h at 1,200 x g at a multiplicity of infection (m.o.i.) of 1 (19). After infection, the cells were washed twice in PBS and cultured in RPMI 1640 media containing penicillin/streptomycin, 10% FCS, and 50 units of IL-2/ml. Supernatant for p24 ELISA was collected 0, 12, 24, and 48 h post-infection. The time points were chosen to allow the virus to finish a full infectious circle (5). CD4+ T cell purity was confirmed by flow cytometry, ranging between 96.3 and 99.8%. The HIV-1JR-FL virus stock was generated by transfection of 293 T cells with pJR-FL and titrated on PBMCs (18).

Cryopreserved PBMC Procedures—Cryopreserved PBMCs were thawed in a 36 °C water bath and were washed twice with PBS at room temperature. 1 x 10^6 cells were lysed immediately, and the remaining cells were positively selected using magnetically labeled CD4 and CD8 microbeads and subsequent column purification according to the manufacturer’s protocol (Miltenyi Biotec). CD4+ T cell purity, verified by flow cytometry, was 97.8% (96.3–99%) (median (range)).

Experimental Design and Statistical Rationale—For the in vitro infection experiment, each time point, including controls, was performed in triplicate. Each sample represents a biological replicate. For the MapDIA statistical model, three biological replicates were sufficient. For the human samples, a total of 10 patients’ samples were obtained. Two patients were excluded due to low cell numbers. Therefore, a total of eight patients were measured at each time point, representing a biological replicate each. We were able to maintain the Replicate Design Model, which enabled us to use the MapDIA statistical model despite the exclusion of two samples.

Proteomics Sample Preparation—Approximately 1 x 10^6 cells were washed with PBS and resuspended in lysis buffer containing 1.2% sodium deoxycholate monohydrate in 50 mM ammonium bicarbonate buffer (150 μl per 1 x 10^6 cells). The cell suspension was thoroughly vortexed and incubated at room temperature for 10 min while shaking at 1,000 rpm. Subsequently, cells were subjected to three 10-min cycles of sonication at 4 °C (100% output, 50% intervals, Branson Sonifier 450, Emerson).

Ice-cold acetone (−20 °C) was added to the lysate at a rate of 6 volumes to 1 volume of lysate and incubated for 1 h on ice. The cell lysate was centrifuged for 15 min at full speed at 4 °C, washed another three times with ice-cold acetone, and then allowed to dry completely. The proteins were resuspended in 8 M urea, 100 mM ammonium bicarbonate buffer.

The protein concentration was estimated using the BCA assay (Pierce). From each sample only 50 μg of protein was used for subsequent steps. Protein disulfide bonds were reduced by adding 5 mM tris(2-carboxyethyl)phosphine and incubating for 30 min at 30 °C. Free cysteine residues were alkylated by adding 40 mM iodoacetamide and incubating for 60 min in the dark at room temperature. Subsequently, the samples were diluted with 0.05 M ammonium bicarbonate buffer to a urea concentration of <2 M, and 1 μg of sequencing grade modified trypsin was added (w/w 1:50). The samples were incubated overnight at 30 °C with shaking at 300 rpm. To stop the tryptic digest, the pH was lowered to 2 using trifluoroacetic acid (final concentration of ~1%) followed by an incubation for 30 min at 37 °C with shaking at 500 rpm.

The cleared peptide solution was desalted with C18 MicroSpin columns (The Nest Group, 5–60 μg loading capacity). Prior to use, the C18 columns were activated with 100% methanol, followed by 80% acetonitrile (ACN), 0.1% TFA, followed by equilibration with 2% ACN, 0.1% TFA. After loading, the columns were washed with 2% ACN, 0.1% TFA. Subsequently, the peptides were eluted with 40% ACN, 0.1% TFA, dried under vacuum, resolubilized in 2% ACN, 0.1% FA, and added at 1:20 with iRT peptides (Biognosys).

Spectral Library and SWATH Peptide Query Parameter Generation—The spectral library and SWATH peptide query parameters were generated essentially as described (20). The TripleTOF 5600 mass spectrometer (AB Sciex) was coupled to a nanoLC 10plus system (Eksigent), and the chromatographic separation of the peptides was performed on a 20-cm emitter (75 μm inner diameter, New Objective) packed in-house with C18 resin (Magic C18 AQ 3 μm diameter, 200 Å pore size, Michrom BioResources). A linear gradient from 2 to 35% solvent B (solvent A, 2% ACN, 0.1% FA; solvent B, 98% ACN, 0.1% FA) was run over 120 min at a flow rate of 300 nl/min. The mass spectrometer was operated in IDA mode with a 500-ms survey scan from which up to 20 ions exceeding 250 counts/s were isolated with a quadrupole resolution of 0.7 Da, using an exclusion window of 20 s. Rolling collision energy was used for fragmentation, and an MS2 spectrum was recorded after an accumulation time of 50 ms. Raw data files (wiff) were centroided and converted into mzML format using the AB Sciex converter (beta version 2011) and subsequently converted into mzXML using openMS (version 1.8). The converted data files were searched using the search engines XTandem (k-score, version 2013.06.15.1) and Comet (version 2013.02, revision 2) against the UniProt-SwissProt complete proteome for Homo sapiens (canonical sequences) with the protein sequences derived from full-length consensus sequences of viral RNA genome of both the HIV-1JR-FL strain used for the in vitro infection model and the viral RNA sequences of 8–10 patients (HIV_fasta_file.txt, supplemental material) (21). The enzyme specificity was set to fully tryptic with two missed cleavages allowed. In total, the database contained 20,149 target sequences and 20,117 decoy se-
sequences. The tolerated mass errors were 50 ppm on MS1 level and 0.1 Da on MS2 level. Carbamidomethylation of cysteines was defined as a fixed modification and methionine oxidation as a variable modification. The search results were processed with PeptideProphet and iProphet as part of the TPP 4.7.0. The spectral library was constructed from the iProphet results with an iProphet probability cutoff of 0.98422, corresponding to a 1% FDR on a protein level as determined by Mayu (version 1.08). The raw and consensus spectral libraries were built with SpectraST (version 5.0) using the -cICID_QTOF option for high resolution and high mass accuracy and -c_IRT and -c_IRR options to normalize all retention times according to the iRT peptides with a linear regression. The six most intense y and b fragment ions of charge state 1 and 2 were extracted from the consensus spectral library using spectrast2tsv.py from msproteomicstools (https://pypi.python.org/pypi/msproteomicstools). Fragment ions falling into the swath window of the precursor were excluded as there is often interference with the resulting signals. Decay transition groups were generated based on shuffled sequences ( decoys similar to targets were excluded) by the OpenMS tool OpenSwathDecoyGenerator (version 1.10.0) and appended to the final SWATH library in TraML format (20).

**SWATH Data Acquisition**—TripleTOF 5600 mass spectrometer was set up as described above but operated in SWATH mode using the following parameters. For liquid chromatography, a linear gradient from 2 to 35% solvent B (98% ACN, 0.1% FA) was run over 120 min at a flow rate of 300 nl/min (patient samples) or 5–30% over 60 min (in vitro infections). Acquisition of a 250-ms survey scan was followed by acquisition of 64 fragment ion spectra from 64 precursor isolation windows (swaths) with variable width chosen to minimize precursor ion density in each window (16). The swaths were overlapping by 1 m/z and covered a range of 400–1200 m/z. The SWATH MS2 spectra were recorded with an accumulation time of 50 ms and cover 100–2000 m/z. The collision energy for each window was determined according to the calculation for a charge 2+ ion centered upon the window with a spread of 15. Raw data files (wiff) were converted into mzXML format using ProteoWizard as described (14).

**SWATH Data Analysis with OpenSWATH**—The SWATH data were analyzed using OpenSWATH version 1.10 with the following parameters. Chromatograms were extracted with 0.05 m/z around the expected mass of the fragment ions and with an extraction window of ±5 min around the expected retention time after iRT alignment. The best model to separate true from false positives (per run) was determined by pyprophet with 10 cross-validation runs. The runs were subsequently aligned using TRIC as described with a target FDR of 0.01 (22).

**Relative Protein Quantification**—To obtain fold changes and corresponding p values of all proteins compared with the time-matched uninfected controls for in vitro infections, or non-viremic patient samples, the software MapDIA was used. Median normalization was applied prior to analysis by MapDIA throughout. A model-based statistical significance analysis of protein level differential expression between the groups of interest was performed as described in the MapDIA package. iRT peptides and serum proteins (samples were stored in BSA containing buffer) were removed from the analysis (supplemental Fig. 1).

**Protein Abundance Estimation**—For label-free abundance estimations of all proteins identified by SWATH MS, the aLFQ package was used (23). Protein abundances were computed as the sum of the top five most intense fragment ions from the three most intense peak groups. The TRIC strategy for automated peptide alignment was used (22). Requantified values were used when no reliable signal was found. A protein had to be detected for a minimum of three times per group in order for the remaining values to be re-quantified and considered for downstream analysis.

**Clustering, Functional Annotation, and Network Analysis**—Clustering was performed using the heatmap2 function in R. For functional annotation, the ClueGO app for Cytoscape was used (24). An Enrichment/Depletion method with a two-sided hypergeometric test was applied, correcting with a Bonferroni step down correction. Min GO level was set to 4 and Max GO level to 10. The protein list from the spectral library was used as background throughout. For networks based on STRING, an evidence grade cutoff of 0.4 was used.

**RESULTS**

**Qualitative Characterization of the HIV-1 and Host Cell Proteomes**—CD4+ T cells from blood donors were isolated and infected in vitro with HIV-1JR-FL (18). The study approach is summarized in Fig. 1A. The HIV-1 protein p24 was quantified by ELISA in the supernatant to confirm successful infection (Fig. 1B). Because human CD4+ T cells need to be activated for successful infection with HIV-1 in vitro, we focused our analysis on the log2 fold changes between paired infected and uninfected cells at a given time point. To generate a spectral library for use as reference in generating peptide query parameters for the peptide-centric analysis of SWATH-MS data, we performed DDA analysis of the proteomes of in vitro HIV-1-infected human CD4+ T cells as well as sorted CD4+ T cells, CD8+ T cells, and PBMC from HIV-1-infected individuals (Fig. 1A and supplemental Fig. 2) following our published protocol (20). We added the sequence of HIV-1JR-FL used for in vitro infection as well as eight full-length HIV-1 sequences from patients to our protein sequence database. This resulted in a reference spectral library (Fig. 1A, qualitative MS), consisting of 3,360 proteins based on 22,901 peptides with an FDR of 1% on the protein level, including four HIV-1 proteins: Env_gp120; Nef; matrix protein (p17); and capsid protein (p24) (Table II, HIV_fasta_file.txt). Using DDA mode for the library creation, we were able to detect HIV-1 proteins from in vitro experiments only, despite adding the matching HIV-1 sequence of the individual virus of each pa-
tient from 8 of the 10 patients. Overall, it is not very surprising that we were unable to find viral proteins in human individuals, given that even in actively replicating patients 1 of ~617–771 of CD4+ T cells is productively infected, i.e. produces viral protein (25). In contrast, after spinoculation with an m.o.i. of 1, close to 100% of CD4+ T cells are infected in vitro.

Changes in the Proteome of in Vitro HIV-1-infected CD4+ T Cells over Time—To determine the changes of the proteome associated with HIV-1 infection, we measured quantitative proteome profiles of CD4+ T cells infected in vitro with HIV-1 by SWATH-MS. Of 1,725 proteins (1% FDR) consistently quantified during the time course, a total of 145 unique proteins (8%) were significantly different between infected and uninfected CD4+ T cells according to a differential protein expression analysis using a Bayesian hierarchical model as implemented in mapDIA (26), using a threshold of (FDR < 0.05 and log2 fold changes >1 and < −1) (supplemental Table 1). The most profound changes in the proteome were detectable 24 h after infection (Fig. 2A). To assess the overall structure of the changes in the proteome, we performed hierarchical clus-

Fig. 1. Experimental design. A, lysates from human CD4+ T cells as well as PBMC, sorted CD4+ T cells, and sorted CD8+ T cells were used to generate a spectral library and HIV-1 sequences (qualitative MS). Using this library, both relative changes between the samples of interest as well as protein abundance (quantitative MS) were calculated using the mapDIA and the aLFQ package, respectively. B, p24 in the cell culture supernatant from infected CD4+ T cells as detected by ELISA.
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| Viral protein | Function | Seq. coverage (%) | distinct peptides |
|--------------|----------|------------------|------------------|
| Env_gp120    | Part of gp160, binds to CXCR4/CCR5 | 7.4              | 3                |
| Nef          | Early protein, inhibits T cell activation | 11.1             | 2                |
| P17          | Part of GAG, structural protein        | 36.9             | 13               |
| P24          | Capsid, forms conical core of virions  | 18.0             | 4                |

Changes in the Proteome of CD4⁺ T Cells from Patients Are Associated with HIV-1 Viremia—Next, we tried to determine whether any of the changes observed in the in vitro HIV-1
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Infection model are recapitulated in CD4+ T cells from HIV-1-infected individuals. Samples were collected from patients participating in the longitudinal Zurich Primary HIV-1 Infection Study. Samples were obtained from patients during viremia as well as at least 6 months after successful antiretroviral treatment where no viral load was detectable (Table I). Therefore, we cannot exclude that the medication itself contributes to the observed changes of the proteome. The clinical samples were heterogeneous in terms of absolute protein yield extracted due to low cell numbers after sorting. Samples with too low amounts of cells were excluded pairwise to retain the Replicate Design Model. We were unable to link to any specific pattern, such as prolonged storage, probably due to the limited sample number. In total, we were able to measure the proteome in two time points each from CD4+ T cells of eight individual patients, and we were able to consistently measure 895 proteins in CD4+ T cells. Next, we compared the quantitative changes in CD4+ T cells in patients during HIV-1 viremia and after successful therapy. Overall, the changes were not very pronounced in terms of log2 fold changes (Fig. 5A). Because CD4+ T cells are the main population of infected cells in peripheral blood, it is to be expected that this cell population shows the biggest changes upon treatment. Using an FDR cutoff of 1% and log2 fold changes of <0.5 to >0.5, as threshold for significance, 57 proteins were up-regulated, and 115 were down-regulated in CD4+ T cells. After functional enrichment analysis, the GO terms RNA polymerase II core promoter, type 1 interferon signaling, neutrophil killing of a symbiont cell, and antigen presentation were enriched in up-regulated proteins (Fig. 5B and supplemental Table 2). Specifically, the small ubiquitin-like protein-modified interferon-stimulated gene 15 (ISG)-15 plays an important role by inhibiting the release of HIV-1 particles (30), and correlates with viral load in humans (31). The rather unexpected finding of increased syntaxin-binding protein (STXBP)-2 is in line with findings in a Jurkat T cell line, showing that STXBP-2 is a strong inhibitor of retroviral transcription (32). RuvB-like 2 (RUVB2), an ATP-dependent DNA helicase, is exploited by HIV-1 to retain the correct ratio of the HIV-1-1 structural protein Gag to the envelope protein Env; in addition, RUVB2 levels correlate with HIV load in humans (33). The detection of the proteins TAP-1 and TAP-2 suggests that there is a contamination of CD4+ T cells with dendritic cells. Dendritic cells

Fig. 2. Changes in the proteome of in vitro HIV-1-infected CD4+ T cells over time. A, violin plots of log2 fold changes of proteins at given time points comparing HIV-1-infected versus -uninfected CD4+ T cells. Single proteins are connected by lines. Box plots show mean ± S.E. An FDR <0.05 and log2 fold changes >1 and <−1 after adjusting for multiple testing were considered significant. The model-based statistical significance analysis from the mapDIA package was used for statistical analysis. B, heat map displaying log2 fold changes of proteins changing at least once significantly (FDR <0.05 and log2 fold changes >1 and <−1) during the time course. Four high level clusters are highlighted in blue (a), green (b), olive (c), and red (d), respectively.
Fig. 3. **Traces and representative single proteins of clusters.** Log2 fold changes of all traces of each cluster and a representative single protein from each cluster are plotted in matching colors to the cluster they represent. Each letter (A–D) represents one of the clusters from Fig. 2. The violin plots represent the distribution of log2 fold change of every protein within the cluster. In the single protein figures, the black dots represent single measurements; the colored dots represent the mean of the log2 fold changes.
are producer of TAP-1 and TAP-2 and are known to express CD4 when isolated from blood from human individuals (34). Hence, a negative sorting approach may be more suitable for this type of analysis. Among the down-regulated proteins, no significant enrichment of GO terms was detectable.

Comparison of in Vitro Infected CD4$^+$ T Cells and CD4$^+$ T Cells Sorted from Patients—Next, we aimed to compare in vitro infected CD4$^+$ T cells and CD4$^+$ T cells from patients. In a first step, we compared how the detected proteome of the two cell populations overlaps (Fig. 7A). 829 proteins were detected both in vivo and in vitro. Then, we compared all proteins considered to be significantly differentially regulated in both groups (Fig. 7B). The overlap was low with only 11 proteins changing both in vivo and in vitro. To assess whether this lack of overlap was due to the fact that the main changes were in proteins that were only detected in one of the groups, we compared all the proteins considered different in the two groups with the total of proteins detected in both groups (overlap, Fig. 7C). In the in vitro experiment, 97 proteins were considered different, which were not detected in the in vivo samples, whereas 37 proteins considered to be regulated in the in vitro time course were detected and considered as
unchanged in the *in vivo* samples. Conversely, in the *in vivo* samples 14 proteins were considered different, which were not detected in the *in vitro* samples, whereas 147 proteins which were differentially regulated in the *in vivo* samples were detected and did not change in the *in vitro* time course. Hence, of the proteins that were detected in both sample types some overlap in differential expression was observed; however, the majority of proteins that were differentially expressed *in vitro* did not change *in vivo* and vice versa.

Functional enrichment analysis showed significant enrichment of type 1 interferon signaling in both *in vitro* infected CD4⁺ T cells as well as CD4⁺ T cells from patients. 4 and 5 proteins *in vitro* and *in vivo*, respectively, were significantly enriched for the GO term “type 1 interferon signaling” (Figs. 4B and 6C). A network based on published interactions using evidence based only on experiments and databases from STRING confirmed that the proteins are interacting based on experimentally observed protein-protein interactions (Fig. 7D). However, the type 1 interferon pathway is a central pathway in innate and adaptive immunity. Furthermore, type 1 interferon signaling may have both protective or detrimental effects for the host, depending on the timing, the state of the cell, and the cellular milieu (35). Consequently, assessing functionality based on the broad GO term “type 1 interferon signaling” might be an oversimplification. Therefore, we plotted the sig-
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Fig. 6. Qualitative and quantitative assessment of the proteome of human CD4+ T cells. A, volcano plot showing log2 fold change of protein abundance plotted against -log10 adjusted p value for CD4+ T cells from viremic patients versus the same viremic patient. Data points highlighted in blue in the upper right section represent proteins that display an enrichment in of log2 fold change >0.5 and adjusted p value <0.05. A total of seven patients was assessed for this comparison. B, network of significantly enriched functional annotation terms of proteins from up-regulated proteins comparing the same patient during viremia and after successful treatment. The GO terms “immune system process” (diamond) and “molecular function” (large circle) were assessed using the GO term “fusion function” from the ClueGO package. Different colors were used for different functional groups. After network analysis of functional terms, the proteins underlying the significantly enriched functional term were added (small circle, protein names are in red). The protein list from the spectral library was used as reference data set. Significance was assessed using a two-sided hypergeometric test with a Bonferroni step down procedure to adjust for multiple testing.

Significantly enriched proteins within a diagram of interferon α/β signaling obtained from the REACTOME database (Fig. 8, A and B) (36). In direct comparison, differentially regulated proteins in vivo affect primarily proteins activating STAT-1, including STAT-1 itself; in turn, differentially regulated proteins in vitro are located downstream of STAT-1 activation in the
pathway, such as mediators of an antiviral state like IRF-4. Hence, despite the significant enrichment of the GO term type 1 interferon signaling \textit{in vivo} and \textit{in vitro}, the actual proteins may be different as well as their biological function (35). In sum, there is little overlap between the specific changes in the proteome comparing \textit{in vivo} and \textit{in vitro} infected CD4$^+$ T cells but rather convergence at the pathway level of type 1 interferon signaling.

**DISCUSSION**

We examined the changes in the proteome of human CD4$^+$ T cells associated with HIV-1 infection both \textit{in vitro} and \textit{in vivo}. The most profound changes in the overall proteome of CD4$^+$ T cells infected with HIV-1 \textit{in vitro} were detected 24 h after infection and involved important functional groups within the cell. In addition, we could measure the abundance and dynamics of certain HIV-1 proteins within the host proteome. Finally, we could reiterate some of the findings from the \textit{in vitro} model using CD4$^+$ T cells from HIV-1-infected individuals.

The proteome of \textit{in vitro} HIV-1-infected human T cells has not been measured yet. Here, we present data from a comprehensive set of samples with longitudinal \textit{in vitro} and \textit{in vivo} samples. In addition, we could detect viral proteins, which
enables us to put them in the context of the host proteome. A strength of our approach is the paired approach for both in vitro and in vivo samples. By comparing cells from the same donor or the same individuals at different time points, we can reduce inter-individual differences.

During the in vitro infection time course, changes in the CD4+ T cell proteome were most prominent 24 h after infection, simultaneous with the peak of viral protein production. GO term enrichment of immune system processes showed a decrease of proteins associated with TLR-4 signaling and type 1 interferon signaling. TLR-4 signaling has been shown to be an important regulator of T cell receptor signaling in T cells from mice (37). Conversely, the HIV-1 envelope protein gp120 has been shown to bind to TLR-4 (38). Our investigation shows a strong time-dependent association of decreased TLR-4 signaling and increased gp120 in the in vitro experiment.

Type 1 interferon signaling plays a central role in viral infections and has pleiotropic effects. Depending on the biological context, type 1 interferons may promote or inhibit T cell

Fig. 8. REACTOME diagrams of the type 1 interferon signaling pathway with highlighted and regulated proteins from the in vivo and in vitro experiments. A, REACTOME diagram of the type 1 interferon signaling pathway highlighted (in purple), with differentially regulated proteins in CD4+ T cells from patients on and off treatment. B, REACTOME diagram of the type 1 interferon signaling pathway with highlighted (in purple) and differentially regulated proteins in CD4+ T cells infected in vitro at the 24-h time point.
activation, proliferation, differentiation, and survival (35). This central role may explain that we could detect changes in proteins associated with type 1 interferons both in vitro and in vivo.

Our enrichment analysis used the protein library only. Therefore, the number of proteins being enriched appears on first sight to be rather low. However, because the number of proteins in our custom protein library for a given GO term is limited, the enrichment is significant even with a low total amount of proteins. For example, four proteins linked to the GO term type 1 interferon changed in the in vitro experiment. The total number of proteins in our library linked to type 1 interferon is 31. Therefore, four changing proteins accounts for ~13% of all type 1 interferon-associated proteins, which is significant.

Our proteome approach and the re-discovery of certain proteins in the context of the whole proteome may be used to support results from genome analyses and put them into biological context. A recent meta-analysis of host cell genes involved in HIV-1 infection emphasized the importance of networks around STAT-1 (39), in line with our in vivo results. Hence, our proteome approach introduces not only an additional level of evidence on the proteome level in vitro but also re-emphasizes and highlights the role of these proteins in actual human individuals.

In addition to host proteins, we could directly detect HIV proteins in the in vitro time course. To the best of our knowledge, we are the first ones to report HIV proteins within the human proteome outside of protein-protein interactions studies (40). We could detect the most abundant late proteins at time point 24 h. The 12-h time may have been too early for the early HIV proteins, such as Tat and Rev (6). The other viral proteins might be expressed at insufficient levels for detection using our single-shot analysis method. This obstacle could be overcome by infecting a more susceptible T cell line that is known to produce much more HIV-1 proteins (5). However, in this study, we focused deliberately on primary CD4+ T cells and clinical samples to assess the translational potential of SWATH-MS. CD4+ T cells produced a significant amount of HIV-1 proteins within 24 h. In this experimental setting, we measure the overall protein production within the CD4+ T cell population, which consists of infected and uninfected T cells. Therefore, from a single cell perspective, the total amount of HIV-1 protein produced is likely to be diluted and very likely to be higher than our estimate, although we tried to minimize the number of uninfected cells by using a multiplicity of infection of 1 in combination with spinoculation to synchronize the infection and enhance the infection rate, i.e. theoretically, each cell should be infected by HIV-1. However, not every primary CD4+ T cell is, even after maximal stimulation, susceptible for HIV-1 infection in vitro (41).

Our results differ to some extent from previous data published on the proteome of in vitro infected T cell lines (7, 42). The study of Chan et al. found changes in 21% of proteins upon HIV infection. However, because T cell lines used are hyper-susceptible for HIV infection compared with human samples, direct comparison to our results is difficult.

Next, we assessed paired CD4+ T cells from patients on and off treatment. Nevertheless, the differences in CD4+ T cells were most pronounced between viremic and non-viremic patients. Whether this is a sampling bias or a real biological phenomenon remains unclear. However, it is tempting to speculate that as CD4+ T cells are the main reservoir of HIV-1 in peripheral blood, they are most prone to changes upon treatment. The number of proteins detected from the clinical samples was low compared with cell lines but was comparable with other clinical tissues (43) and similar to a previous study that analyzed PBMCs by SWATH-MS (44). As such, optimization of the sample preparation from patient-derived PBMCs, combined with a deeper coverage in the spectral library via fractionation, may yield improved proteome coverage in future studies. A further limitation of the in vivo component of our study was the relatively limited numbers of patient samples analyzed, which resulted in a relatively low number of proteins called as differentially expressed likely reflecting the large differences in genetic and environmental backgrounds of the patients. Hence, because of the heterogeneity of data, the patient samples represent a pilot analysis, and a larger number of samples is required for more statistical power to overcome strongly confounding signals not related to the viral infection.

Even though we used primary human CD4+ T cells for the in vitro infection, it is difficult to compare these findings directly to the changes detected in the proteome of patients during HIV-1 infection. First and foremost, the relationship of infected versus bystander cells is fundamentally different. In vitro, we expect a high percentage of infected cells, whereas in vivo the overwhelming majority of cells is not infected (41). In addition, for an in vitro infection of human CD4+ T cells to be successful, the cells must be artificially activated. Hence, the in vitro data in our investigation display the changes within the proteome against a strong background of activation. We tried to be as specific as possible to detect only changes associated with HIV-1 by comparing activated and simultaneously HIV-1-infected cells against activated cells only at the same time points. This approach notwithstanding, the detectable changes are still against a background of high activation. In humans, in contrast, inflammation is associated with viremia; suppression of viremia, in turn, abrogates inflammation. Hence, the reported comparison of T cells from viremic versus non-viremic patients includes changes attributable to resolving inflammation as well. Furthermore, the changes detectable over time in the in vitro system with a defined infection time point are likely to occur at the same time in an infected individual, for example, infection, integration, and assembly of the virus are likely to happen constantly in humans and successively in vitro. Finally, differences in the host (such as genetic differences, differences within the virus, and differences in the handling of clinical samples) contribute to
increase the noise within the measurements. Despite all these drawbacks, we could link changes in the proteome both in vivo and in vitro, for example with proteins associated with type 1 interferon signaling. However, it is worth noting that there may be relevant biological differences in the differentially expressed proteins associated with the GO term “type 1 interferon signaling” in the in vivo and the in vitro samples.

In sum, we analyzed the changes of the proteome of human CD4+ T cells associated with HIV-1 infection in both an in vitro model of infection and cells derived from HIV-1-infected patients. To our best knowledge, the proteome of CD4+ T cells from HIV-infected humans has not been measured before. Our results warrant further studies with larger patient numbers and different clinical end points to better understand the host immune system in HIV-1 infection.

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DATA AVAILABILITY

The mass spectrometry proteomics data has been deposited to the ProteomeXchange Consortium (http://proteomecentral.proteomexchange.org) via the PRIDE partner repository (45) with the data set identifier PXD005234.

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