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Lapatinib-Binding Protein Kinases in the African Trypanosome: Identification of Cellular Targets for Kinase-Directed Chemical Scaffolds

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Abstract

Human African trypanosomiasis is caused by the eukaryotic microbe Trypanosoma brucei. To discover new drugs against the disease, one may use drugs in the clinic for other indications whose chemical scaffolds can be optimized via a medicinal chemistry campaign to achieve greater potency against the trypanosome. Towards this goal, we tested inhibitors of human EGFR and/or VEGFR as possible anti-trypanosome compounds. The 4-anilinoquinazolines canertinib and lapatinib, and the pyrrolopyrimidine AEE788 killed bloodstream T. brucei in vitro with GI50 in the low micromolar range. Curiously, the genome of T. brucei does not encode EGFR or VEGFR, indicating that the drugs recognize alternate proteins. To discover these novel targets, a trypanosome lysate was adsorbed to an ATP-sepharose matrix and washed with a high salt solution followed by nicotinamide adenine dinucleotide (NAD³). Proteins that remained bound to the column were eluted with drugs, and identified by mass spectrometry/bioinformatics. Lapatinib bound to Tb927.4.5180 (termed TbLBPK1) while AEE788 bound Tb927.5.800 (TbLBPK2). When the NAD³ wash was omitted from the protocol, AEE788, canertinib and lapatinib eluted TbLBPK1, TbLBPK2, and Tb927.3.1570 (TbLBPK3). In addition, both canertinib and lapatinib eluted Tb10.60.3140 (TbLBPK4), whereas only canertinib desorbed Tb10.61.1880 (TbCBPK1). Lapatinib binds to a unique conformation of protein kinases. To gain insight into the structural basis for lapatinib interaction with TbLBPKs, we constructed three-dimensional models of lapatinib-TbLBPK complexes, which confirmed that TbLBPKs can adopt lapatinib-compatible conformations. Further, lapatinib, AEE788, and canertinib were docked to TbLBPKs with favorable scores. Our studies (a) present novel targets of kinase-directed drugs in the trypanosome, and (b) offer the 4-anilinoquinazoline and pyrrolopyrimidines as scaffolds worthy of medicinal chemistry and structural biology campaigns to develop them into anti-trypanosome drugs.

Introduction

Human African trypanosomiasis (HAT) is caused by the protozoan Trypanosoma brucei. New drugs for the treatment of HAT are needed as the current ones have significant issues relating to administration route, toxicity and/or drug resistance [1,2]. To assist drug discovery efforts, we must find novel small molecules that kill the parasite, and, in the long-term, also understand the mechanisms of their action in the trypanosome.

Protein phosphorylation on Tyr residues, catalyzed by protein Tyr kinases (PTKs) or dual-specificity protein kinases that act on Tyr as well as Ser/Thr [3–6], is important for cell signaling networks that control numerous processes [7]. PTKs in vertebrates exist in two general forms; receptor tyrosine kinases (RTKs) and non-receptor Tyr kinases (NRTKs). RTKs such as epidermal growth factor receptor (EGFR) or vascular endothelial growth factor receptor (VEGFR), are transmembrane proteins and are activated upon binding of hormones to their extracellular domains. NRTKs, e.g., Src-family protein kinases, are cytosolic, and can be activated by RTKs or cytokine receptors.

Mechanistic studies of phosphotyrosine (pTyr) signaling in human cells led to discovery of drugs (e.g., lapatinib, and imatinib) that are incredibly well-tolerated [8–10]. Lapatinib (GW572016) (GlxoSmithKline) inhibits human EGFR/HER2 [11]. AEE788 (Novartis) inhibits EGFR and VEGFR [12,13], and canertinib (CI-1033) (Pfizer) is a pan-inhibitor of EGFRs [14–16].

T. brucei has a divergent pTyr system, as compared to its human host. Whereas Tyr-phosphorylated proteins are present [17,18], trypanosomes lack RTKs [19,20]. Consequently, Tyr-phosphorylation of proteins is projected to be performed by dual specificity enzymes that act on Ser/Thr as well as Tyr residues [19]. Further, the trypanosome genome does not encode classic pTyr-binding domains (e.g., SH2) [21]. The biological importance of Tyr-phosphorylation in T. brucei is inferred from the effect of small molecule Tyr kinase inhibitors; genistein blocks trypanosome...
replication [22]; (ii) Tyrphostin A47 inhibits interferon-γ induced Tyr phosphorylation of proteins [23], and (iii) Tyrphostin A47 inhibits receptor mediated endocytosis by the parasite [24,25].

Due to the uniqueness of the Tyr-phosphorylation system in the trypanosome it seems likely that a detailed understanding of pTyr signaling (i.e., enzymes, substrates and pathways regulated) could yield great insight into novel signal transduction pathways in a deeply-diverged eukaryote, and lead to discovery of novel targets for lead drug discovery.

In the work reported here, we show that the 4-anilinoquinazolines lapatinib and canertinib kill bloodstream T. brucei. Since the trypanosome genome does not encode orthologs of EGFR/HER2, we searched for and identified novel trypanosome targets of the drugs, using an affinity chromatography protocol that involved protein elution with drugs: Proteins were identified with mass spectrometry and bioinformatics. This protocol which is easily adaptable for discovering targets of kinase inhibitors in other cells, led to identification five novel binding-proteins of lapatinib, AEE788, and canertinib in T. brucei. The four putative targets of lapatinib were evaluated theoretically for their ability to bind the drug using homology modeling and molecular docking. Results from these studies support conclusions drawn from our ligand-affinity chromatography data.

Materials and Methods

Reagents

Immobilized γ-aminophenyl-ATP-sepharose was purchased from Jena Biosciences. NAD+, and ATP were purchased from Sigma. Silver stain kit was obtained from Pierce. Lapatinib, canertinib, and AEE788 were gifts from GlaxoSmithKline, Pfizer, and Novartis, respectively.

Effects of Drugs on Trypanosome and HeLa Cell Replication

Bloodstream form T. brucei Lister 427 were seeded at a density of 2×10^5 cells/ml and cultured in HMI-9 medium [26] in a 24 well plate. Two microliters of DMSO (control) or different concentrations of drugs (as specified in the Figure legends) from 200X DMSO stocks were added to the cultures. Cells were inoculated at 37°C for 48 h, and counted with a haemocytometer.

HeLa cells were cultured in high glucose Dulbecco’s modified Eagle’s medium (Cellgro) supplemented with 10% FBS (HyClone), 100 U/ml penicillin (Cellgro), 100 µg/ml streptomycin (Cellgro) [27]. Cells were cultured at 37°C with humidified 5% CO2. For drug inhibition studies, cells were seeded in a 24 well plates at a density of 1×10^5 cells/ml. Drugs or DMSO were added in 2 µl up to 25 µM final concentration. Cells were incubated at 37°C for 48 h and counted with a haemocytometer.

Drug Affinity Chromatography of Proteins

T. brucei were cultured axenically [26] to a density of 1×10^6 cells/ml and harvested (2×10^6 total cells). After washing with 10 ml of cold BBS/G (buffered saline plus glucose; 50 mM bicine, 150 mM NaCl, 5 mM KCl, 1% glucose, pH 7.4) containing 1 mM sodium vanadate, cells were lysed on ice in 1 ml of cold Buffer A [20 mM Tris-HCl pH 7.4, 60 mM MgCl2, 60 mM KCl, 1 mM DTT, 0.2% NP-40, 1 mM PMSF, 2 mM L-azophenylalanine, 5 µg/ml leupeptin, 10 µM TLCK, 2 µM FMK024, and 1 mM sodium vanadate]. The cell lysate was incubated on ice for 20 minutes and centrifuged at 15,000×g for 10 min at 4°C. The supernatant was incubated for 18 h at 4°C with 50 µl (settled bed volume) of ATP-sepharose (Sepharose 4B was used as control in a separate experiment.) The resin was washed with 500 µl each of the following (i) buffer A; (ii) buffer A containing 1 M KCl, and finally (iii) buffer A. When specified, the resin was eluted twice with 100 µM NAD+ (100 µM in buffer A).

Several concentrations of drugs (i.e., 1 µM, 10 µM, and 100 µM) were used in trial experiments to elute proteins from the ATP-affinity column. For (subsequent) mass spectrometry analysis it is important that the bands detected by silver staining of protein be sufficiently dark. After analysis of the proteins eluted in several independent trial studies, we determined that 100 µM concentration was optimal for elution of proteins bound by lapatinib, AEE788 and canertinib: The drug concentration needed varies for each scaffold. For each drug, proteins bound to the affinity column were eluted twice (from different matrices) with 100 µl of lapatinib (100 µM) or AEE788 (100 µM) or canertinib (100 µM) (Drugs are diluted to their final concentrations in buffer A from DMSO stocks at 10 mM concentration). A control elution was performed with buffer A containing 1% DMSO, because the final concentration of that solvent in the drug mixtures was 1%, and eluted proteins identified by mass spectrometry. Eluted polypeptides were precipitated with trichloroacetic acid and pelleted (15,000×g for 10 min). The supernatant was aspirated, the precipitate was suspended in 2 µl of water, the supernatant was diluted to 100 µl, and the pH adjusted by adding 5–7 µl of 30% NH4OH. Proteins were separated by SDS-PAGE (12%), and polypeptide bands detected by staining with silver. Each lane of gel was cut into four pieces each of which was mixed with a sterile blade in preparation for trypsinization (see below).

In-gel Trypsin Digestion and Mass Spectrometry

Polyacrylamide gel pieces were destained with a kit (Invitrogen), and dehydrated with acetonitrile. Proteins were digested overnight with trypsin (5 ng/µl Promega) in ammonium bicarbonate (50 mM) at 37°C. Peptides were extracted with formic acid (5% v/v in water) for 30 min, and then with acetonitrile. Extracts were pooled, dried in a SpeedVac, and peptides were purified with ZipTip™ C18 chromatography (Millipore Corporation).
**Table 1.** Protein kinases eluted with drugs after 1 M KCl wash of the affinity column.

| Lapatinib               | Canertinib            |
|-------------------------|-----------------------|
| **Protein**             | **Peptides**          | **Peptide Sequence** | **Residue No.** |
| Tb927.4.5180 (TbLBPK1)  | 6                     | K.QQQQDLNHEK.K      | 280–291         |
|                         |                       | R.DEVEELK.K         | 220–229         |
|                         |                       | K.AFDLQEAR.Y        | 336–345         |
|                         |                       | R.RPFAEGESQQQWQNL.K | 537–554         |
|                         |                       | R.QLTMQLEESVR.R     | 208–221         |
|                         |                       | K.Q.ATLPSYGLNATVFR.K | 92–110     |
| Tb927.5.800 (TbLBPK2)   | 4                     | K.TRHPQLAFAE.R      | 45–57           |
|                         |                       | R.GTNIOTGDQPAIK.L  | 27–41           |
|                         |                       | K.TLMLAEQMIA.R     | 108–121         |
|                         |                       | R.GSLLPWQLK.A      | 213–223         |
| Tb927.3.1570 (TbLBPK3)  | 2                     | R.RPGLCDSPSLEAF.K   | 118–133         |
|                         |                       | K.ASLFTDLPTAATLPK.R | 497–514         |
| Tb10.61.3140 (TbLBPK4)  | 2                     | R.VAGQGTFTGVQLARD  | 24–39           |
|                         |                       | K.Q.QLPAEVYDLCGK.I | 276–290         |

**LC-MS/MS analysis** was performed with a LTQ mass spectrometer (Thermo Scientific). The LC system was configured in a vented format consisting of a fused-silica nanospray needle packed in-house with Magic C18 AQ 100 A reverse-phase media (Michrom Bioresources Inc.), and a trap containing Magic C18 AQ 200 A [30]. Peptide samples were loaded onto the column and separated using a two-mobile-phase system consisting of 0.1% formic acid in water (A) and 0.1% acetic acid in acetonitrile (B). The mass spectrometer was operated in a data-dependent MS/MS mode over the m/z range of 400–1800. For each cycle, the five most abundant ions from the scan were selected for MS/MS analysis using 35% normalized collision energy. Selected ions were dynamically excluded for 45 seconds. Raw MS/MS data were submitted to the Computational Proteomics Analysis System (CPAS), a web-based system built on the LabKey Server v11.2 that utilizes Trans-Proteomic Pipeline v4.3 [28,29], and searched using X! Tandem engine (2009.10.01.1) against Trypanosome protein database v. 4.0 (ftp://ftp.sanger.ac.uk/pub/databases/T. brucei_sequences/T.brucei_genome_v4/), which included additional common contaminants such as human keratin. The following modifications were considered: loss of water or ammonia from terminal glutamic acid or glutamine respectively, and oxidation of methionine. The mass tolerances were set ±2 Da and ±0.5 Da for precursor and fragment ions respectively. The enzyme was set to Trypsin, and up to 2 missed cleavages were permitted. The search output files were analyzed and validated by ProteinProphet [30,31]. Peptides matched with an error rate <0.05 were accepted, and corresponding proteins identified with two or more peptides are reported. Experiments were repeated thrice; proteins are reported only if they were detected in at least two separate experiments (Table 1). Properties of peptides used to identify protein kinases are presented in Table S1.

**Structure-directed Sequence Alignment of Lapatinib-binding Pockets**

All four trypanosome kinases have medium to high homology templates in PDB (Table 2). The closest homology templates for TbLBPK1 and TbLBPK3 in the PDB are a putative calcium-dependent protein kinase 1 from Cryptosporidium parvum (CpCDPK1, Uniprot ID A3FLQ16, 34% sequence identity to TbLBPK1 in the kinase domain) and a putative calcium-dependent protein kinase 3 from Toxoplasma gondii (TgCDPK3, Uniprot ID Q3HNM6, 30% sequence identity to TbLBPK3 in the kinase domain). Despite belonging to CaMK family of Ser/Thr kinases, both of these templates are crystallized in lapatinib-compatible backbone conformations (PDB codes 2wei and 3hzt, respectively). In the case of TbLBPK2 and TbLBPK4, the closest homology templates human casein kinase Iβ (Uniprot ID KC1D_HUMAN) and a putative glycogen synthase kinase from Leishmania major (Uniprot ID Q4Q155), are crystalized in an active-like, and therefore lapatinib-incompatible conformation. Thus, the closest template models of TbLBPK1 and TbLBPK3 were capable of favorably docking lapatinib while the closest template for TbLBPK2 and TbLBPK4 could not. We hasten to add that the lapatinib-conformation is dynamic; it is an alternate to the enzyme active-conformation. Therefore, its absence from an x-ray structure does not preclude its existence; in some instances presence of the drug might stabilize lapatinib-compatible conformations in select protein kinases.

Given this background (above), we enhanced modeling based on closest homology templates with more distant modeling protocols using structures of human EGFR in complex with lapatinib (PDB 1skkA [32]) as templates, since our affinity chromatography studies identified TbLBPKs as lapatinib-binding polypeptides. These structures provide...
binding was discerned. From these, lapatinib-propagation of the obtained residue correspondence onto the target closest homology templates of TbLBPKs in PDB (Table 1) followed by ambiguities by constructing structural alignments of EGFR to the several regions of the kinase domain. We resolved the alignment methods failed to find the correct residue correspondence in reliable homology modeling [33–35]. Consequently, sequence align-
tions between EGFR and TbLBPKs is between 18% and 22%, which is below the 40% cutoff for modeling studies. Sequence identity between EGFR and ed a variety of suitable backbone conformations of the kinase domains for the modeling studies. Sequence identity between EGFR and TbLBPKs is between 18% and 22%, which is below the 40% cutoff for reliable homology modeling [33–35]. Consequently, sequence alignment methods failed to find the correct residue correspondence in several regions of the kinase domain. We resolved the alignment ambiguities by constructing structural alignments of EGFR to the closest homology templates of TbLBPKs in PDB (Table 1) followed by propagation of the obtained residue correspondence onto the target sequences via a sequence-based alignment. From these, lapatinib-binding was discerned.

Homology Modeling of TbLBPK Structures

The structures of TbLBPK1-4 and TbCBPK1 were built using the homology modeling platform of the Internal Coordinate Mechanics (ICM) software [36–38]. Initially, well-aligned regions of the target kinase were threaded through the backbone coordinates of the template structure. Loops, insertions and deletions were searched against a large database of PDB fragments for similar sequence and termini topology; well-scoring fragments were incorporated into the nascent model and minimized in its context. Template hydrogen bonds were converted into distance restraints; the model side-chains were thoroughly sampled to find the global minimum of the energy function that included soft van der Waals, electrostatic, hydrogen bonding, torsional strain, and distance restraint terms. To resolve the remaining steric conflicts, the model was subjected to gradient minimization with both side-chain and backbone variables relaxed. The final models were evaluated using two methods: first, ICM Protein Health evaluation confirmed the absence of torsional or steric inaccuracies; second, PROCHECK [39] evaluation placed at least 80% of the residues in the most favorable regions on the Ramachandran plot with at least 97% being in the allowed regions. Visual inspection of the remaining (unfavorable) residues confirmed that they located in extended loops distant from the lapatinib binding site and were therefore not influential in the docking studies.

Ligand Docking and Scoring

ICM ligand docking is based on biased probability Monte Carlo optimization of the ligand internal coordinates in the set of grid potential maps that represent the protein binding pocket. A diverse set of ligand conformers was first generated from 2D coordinates by 2D to 3D conversion and thorough sampling in vacuo. The conformers were placed into the binding pocket in four principal orientations and used as starting points for Monte Carlo optimization. The optimized energy function included the ligand internal strain and a weighted sum of the as starting points for Monte Carlo optimization. The optimized energy potential maps that represent the protein binding pocket. A diverse set of ligand conformers was first generated from 2D coordinates by 2D to 3D conversion and thorough sampling in vacuo. The conformers were placed into the binding pocket in four principal orientations and used as starting points for Monte Carlo optimization. The optimized energy function included the ligand internal strain and a weighted sum of the

S_{bind} = E_{tot} + T \Delta \Delta S_{Tot} + E_{non} + a_1 E_{vdW} + a_2 E_{elec} + a_3 E_{nbs} + a_4 E_{st}

where $E_{tot}$, $E_{non}$, $E_{vdW}$, $E_{elec}$, and $E_{st}$ are Van der Waals, electrostatic, hydrogen bonding, non-polar and polar atom solvation energy differences between bound and unbound states, $E_{vdW}$ is the ligand internal strain, $\Delta \Delta S_{Tot}$ is its conformational entropy loss upon binding, $T = 300$ K, and $z_i$ are ligand- and protein-independent

### Table 2. TbLBPKs1-4 and their structural homologs in PDB.

| Protein Kinase | System ID | UniProt ID | Kinase Domain | Closest PDB SwissProt ID | homologs Sequence Identity to Target (%) |
|---------------|-----------|------------|---------------|--------------------------|-----------------------------------------|
| TbLBPK1       | Tb927.4.5180 | Q58418 | 320:601 | SNF1_YEAST, A3FQ16_CRYPV | 31, 34 |
| TbLBPK2       | Tb927.5.800  | Q57W25  | 11:282 | KC1D_HUMAN               | 69 |
| TbLBPK3       | Tb927.3.1570 | Q57X20  | 213:488 | MARK2_HUMAN, B6KR85_TOXGO | 32, 30 |
| TbLBPK4       | Tb10.61.3140 | Q388M1 (GSK3β) | 20:310 | Q4QE15_LEIMA, GSK3B_HUMAN | 69, 54 |
| TbCBPK1       | Tb10.61.1880 | Q388C6  | 16:284 | M3K5_HUMAN, PAK1_HUMAN | 37.5, 37.3 |

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constants that have been previously optimized on a diverse screening benchmark.

Results

Lapatinib, Canertinib and AEE788 Inhibit Replication of T. brucei at Low Micromolar Concentrations

In the African trypanosome, endocytosis of transferrin, which is necessary for replication [41], is inhibited by Tyrophostin A47, a pan-Tyr kinase inhibitor that also kills the parasite [24]. This data suggests that chemical scaffolds of drugs that inhibit protein Tyr kinases have potential to be “re-purposed” for anti-trypanosome lead drug discovery.

Lapatinib, canertinib and AEE788 (Fig. 1) are Tyr kinase inhibitors that have passed phase I clinical trials, at least, for use in humans [42–45]. In addition, the chemistry for synthesis of these drugs is well established and streamlined for optimization [11,46,47]. To evaluate effects of these drugs on T. brucei, axenically cultured bloodstream cells were exposed to different amounts of the compounds, and checked for viability. Lapatinib killed T. brucei (GI50 = 1.5 μM (Fig. 2A)). Similarly, canertinib and AEE788 killed the trypanosome with an GI50’s of 2 μM, and 3 μM, respectively (data not presented). Ability of lapatinib to interfere with pro-survival pathways (i.e., kill within the half-time of cell replication) in T. brucei was tested directly. When cells (106/ml) were incubated with the drug the survival half-life was 3 h (Fig. 2C), which is shorter than the cell division time of 6–8 h.

Toxicity of lapatinib against T. brucei was compared to its effect on a human HeLa cell line. At 5 μM or 10 μM all trypanosomes were killed but the human cells survived (Fig. 2B), presumably because growth of HeLa is not dependent on a hyperactive EGFR/HER2 [9,12]. Selectivity of lapatinib against the trypanosome is similar to that observed for the drug when tumor cells are compared to untransformed (non-sensitive) human cells [11,48].

Lapatinib, Canertinib and AEE788 Bind Five Trypanosome Protein Kinases

The respectable whole cell potency (i.e., in a phenotypic screen) of lapatinib, canertinib and AEE788 against T. brucei (Fig. 2) could not be explained by inhibition of EGFR and/or VEGFR kinases, because the trypanosome genome does not encode classic receptor Tyr kinases [49]. Therefore, we postulated that the drugs had alternate protein targets in the trypanosome.

To identify the trypanosome targets of the drugs we used them as chemical tools. In our approach, a cell lysate from T. brucei was adsorbed to an ATP-sepharose column. After extensive washing of the ATP-affinity column with buffers containing high salt (i.e., 1 M KCl) and/or NAD+ (100 μM), the drugs (100 μM) were used to elute proteins remaining on the column. Proteins eluted were identified by mass spectrometry and bioinformatic analysis [29,50,51]. The profile of proteins eluted with drugs after the high salt wash is shown in Fig. 3B–D, and protein kinases identified are listed in Table 1. In preliminary studies, the profile of eluted proteins was very similar for the 1 or 10 or 100 μM drug concentrations. The higher concentration of drug eluted more proteins from the affinity column, judging from the intensity of the silver-stained bands and the number proteins identified by mass spectrometry (see Table S2 and Table S3); it was used for our studies.

Only five protein kinases, out of a possible 186 in the trypanosome genome [19,52] bound the three drugs. Tb927.3.1570, Tb927.4.1580 and Tb927.5.800 were eluted with AEE788, canertinib and lapatinib (Table 1). Tb10.60.3140 was eluted by both canertinib and lapatinib, whereas Tb10.61.1880 was only eluted by canertinib. Three lapatinib-binding protein kinases (TbLBPKs) have been studied earlier; TbLBPK1 (termed a Tousled-like kinase) [53,54], TbLBPK-2 (TbCK1.2 [55]) and TbLBPK4 (TbGSK3β [56]). Non-kinase ATP-binding proteins detected in our dataset are available in Table S2. We focused primarily on protein kinases because lapatinib binds that class of enzymes in human cells [57].

Inclusion of NAD+ in Washes of the Affinity Column Alters Profile of Protein Kinases Eluted by Drugs

Cells contain purine cofactors and nucleotides, including NAD+, whose structure overlaps significantly that of ATP, which is a substrate of protein kinases. Recognizing the possibility that
protein kinases may bind weakly to NAD$^+$, we explored a second strategy for our chemical proteomics study. Here, after washing the affinity column with the high salt buffer, we eluted the matrix with NAD$^+$ (100 μM) before desorbing the residual proteins with drugs (see Fig. 3). Protein kinases in drug eluates are presented in Table 3.

After the NAD$^+$ wash, only two protein kinases were eluted from the affinity column. Tb927.4.5180 was eluted with lapatinib, and Tb927.5.800 was found in the AEE788 eluate. Canertinib failed to elute any protein kinase with this revised protocol. One explanation for these data, compared to proteins protein kinases, and found Tb927.5.800, Tb10.70.2070, Tb10.61.1880, and Tb10.61.3140. We conclude that NAD$^+$ can elute select trypanosome protein kinases from an ATP-affinity column, and alter the profile of proteins eluted by drugs from the ATP-affinity matrix.

Binding of Lapatinib or NAD$^+$ to TbLBPKs is Affirmed by Molecular Modeling

The affinity chromatography data indicates that lapatinib, as a model inhibitor, binds to 4 protein kinases (Table 1). Yet an NAD$^+$ wash of the column left Tb927.4.5180 was eluted with lapatinib, and Tb927.5.800 was found in the AEE788 eluate. Canertinib failed to elute any protein kinase with this revised protocol. One explanation for these data, compared to proteins eluted without the NAD$^+$ wash (compare Table 1 and Table 3), is that some protein kinases were eluted by NAD$^+$. We evaluated this hypothesis by analyzing the NAD$^+$ eluates for protein kinases, and found Tb927.5.800, Tb10.70.2070, Tb10.61.1880, and Tb10.61.3140. We conclude that NAD$^+$ can elute select trypanosome protein kinases from an ATP-affinity column, and alter the profile of proteins eluted by drugs from the ATP-affinity matrix.

Lapatinib-Binding Proteins from a Trypanosome

**Figure 3. Drug elution of T. brucei protein kinases using an ATP-affinity resin.** Proteins in total cell lysate from T. brucei were bound on ATP resin. Sepharose 4B resin was used as control. Unbound proteins (Flow through) were recovered and resin was washed sequentially with buffer A and buffer A containing 1 M KCl (Panel A). Bound proteins were eluted with lapatinib (100 μM), or AEE788 (100 μM) or canertinib (100 μM) (Panel B, C and D, respectively). Proteins were visualized by silver staining. Matrix label, A is ATP-sepharose, and S is Sepharose 4B.

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| Protein   | Description               | Lapatinib | AEE788 | Canertinib |
|-----------|----------------------------|-----------|--------|------------|
|           |                           | Total peptides | Total peptides | Total peptides |
| Tb927.4.5180 | Protein Kinase putative   | 6         | 0      | 0          |
| Tb927.5.800  | Protein Kinase             | 0         | 3      | 0          |

Proteins listed are detected at least twice in three separate affinity chromatography/mass spectrometry studies.
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The models constructed as described in Materials and Methods helped to elucidate possible structural basis of TbLBPK interaction with lapatinib and AEE788 and to identify amino residues that shape the drug-binding pocket (Fig. 5). This computational model confirmed that TbLBPKs can adopt the lapatinib-compatible backbone conformation, and that in all four cases, pocket residues are positioned favorably for lapatinib binding (Fig. 5A).

Docking scores for different kinase-drug complexes were consistent with the affinity chromatography data (Table 1 and Table 3): lapatinib, AEE788, and canertinib scored most favorably in the models of TbLBPK1, TbLBPK2, and TbCBPK1, respectively (Fig. 6A). The best poses for the drug binding to the protein kinases are depicted in Fig. 6B. In comparing the chromatography data with the models we found than an arbitrarily chosen ICM score cutoff of -42 separates drug/kinases complexes that are resistant to NAD$^+$ elution from the ATP-affinity column. For canertinib, the results obtained make for interesting discussion: TbCBPK1 docks best to canertinib (among the three drugs tested). However the ICM score was less than 42 for all three drugs, suggesting that TbCBPK1 will not be eluted by any drug from the affinity column after an NAD$^+$ wash (Fig. 6A). These data suggest that the models could have predictive value: a ligand-TbLBPK complex that receives a score $\geq$41 or less is not likely to be eluted from the ATP-affinity column after an NAD$^+$ wash (Table 3 and Table 4).

How could one explain the decreased docking scores, as compared to TbLBPK1, for TbLBPK2 and TbLBPK4? Here the

Figure 4. Closest homology templates for TbLBPK1 (magenta) and TbLBPK3 (navy) are crystallographically observed in a conformation similar to that of lapatinib-bound human EGFR (white).
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Figure 5. Homology modeling of T. brucei lapatinib-binding kinases using structures of human EGFR as templates. (A) Alignment of lapatinib binding site residues. (B) Binding site structures colored according to the alignment. (C) Models of TbLBPK's1-4 complexed with lapatinib.
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decrease in predicted binding affinity of lapatinib may be explained by a missing residue in the hinge region (as compared to human EGFR), which leads to non-optimal orientation of the backbone amide groups. In the case of TbLBPK3, the likely reason for a docking score decrease is a modified environment for the fluorophenyl moiety of the drug, which, for example, includes a non-conservative substitution of Leu in EGFR for Tyr in TbLBPK3 (β4 strand of the kinase) (Fig. 5).

We also tested whether TbLBPKs bind NAD+ as postulated earlier. For this objective, we constructed models of TbLBPKs in the ATP-compatible conformation and docked ATP or NAD+ into the models. Due to high structural similarity between ATP and NAD+, both molecules fit well in the binding pockets and make similar hydrogen bonds with the hinge region and with the conserved Lys in the β3 strand of the kinase. However, entropic and solvation electrostics penalties for the larger and more flexible NAD+ molecule led to better binding scores for ATP (Fig. 7). These results indicate that high concentrations of NAD+ are likely to compete with ATP for binding to TbLBPK and may wash some TbLBPKs off the ATP-affinity column. This prediction is confirmed by detection of Tb927.10.5140, Tb927.5.800 (TbLBPK2), and Tb10.61.1880 in the NAD+ eluate from the affinity chromatography.

**Table 4.** An NAD+(+) wash of affinity columns alters ligand selection of target proteins.

| Protein     | Lapatinib | AEE788 | Canertinib |
|-------------|-----------|--------|------------|
|             | KCL       | KCI/NAD+ | KCL      | KCI/NAD+ | KCL       | KCI/NAD+ |
| Tb927.3.1570  | +         | −       | +         | −         | +         | −         |
| Tb927.4.5180  | +         | +       | +         | −         | +         | −         |
| Tb927.5.800   | +         | −       | +         | +         | +         | −         |
| Tb10.61.1880  | −         | −       | −         | +         | −         | +         |
| Tb10.61.3140  | +         | −       | −         | +         | −         | +         |

Figure 6. Best models of TbLBPK-drug complexes are consistent with affinity chromatography elution data. (A) Predicted ICM binding scores of lapatinib (red), AEE788 (green) and canertinib (magenta) in the binding pockets of the protein kinases. The dotted line represents a hypothetical cutoff for kinase elution by drug after an NAD (+) wash of the affinity column. (B) Predicted binding poses for lapatinib (red), AEE788 (green) and canertinib (magenta) to their highest affinity protein kinases; TbLBPK1, TbLBPK2, and TbCBPK1, respectively. Kinase hinge region is shown in ribbon, ligand atom placement surface is represented by a wire mesh and colored according to its binding properties.

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ATP-affinity column (not presented). Interestingly, one protein kinase eluted by NAD\(^+\), i.e., Tb927.10.5140 was not eluted by any of the drugs used in this study.

Discussion

New Chemical Entities for Anti-Trypanosome for Hit-to-Lead Studies

Current anti-trypanosome chemotherapies are difficult to deliver safely and some can be very toxic [1]. For example, in late stage human African trypanosomiasis (HAT) when the parasite has entered the central nervous system, the arsenical drug melarsoprol is the frontline treatment despite its toxicity (5% of patients die from side effects). Consequently, a pressing need exists for new therapeutics with oral bioavailability and good safety profile.

We discovered that endocytosis of transferrin by bloodstream \(T. brucei\) is inhibited by Tyrphostin A47 [24,25], a pan-inhibitor of protein Tyr kinases that also kills the parasite (Subramanya and Mensa-Wilmot, unpublished). We inferred from these data that inhibitors of \(T. brucei\) protein Tyr kinases or dual-specificity protein kinases (trypanosomes lack classic Tyr kinases (e.g., EGFR)) might be valuable as “hits” for anti-trypanosome lead drug discovery. After performing a “focused screen” of investigational or approved Tyr kinase drugs, we settled on the 4-anilinoquinazolines canertinib (CI-1033) [14,46,61] and lapatinib (GW572016) [42,58], and the pyrrolopyrimidine AEE788 [12,62] (Fig. 1) as possible “hits” for anti-trypanosome lead discovery. Canertinib, lapatinib, and AEE788 killed bloodstream \(T. brucei\), with GI\(_{50}\) of 3 \(\mu\)M or less (Fig. 2).

The biological importance of three \(TbLBPK\)s has been characterized in RNA interference studies; \(TbLBPK1\) (previously named Tousled-like kinase) [53,54], \(TbLBPK2\) (TbCK1.2 [55]) and \(TbLBPK4\) (TbGSK3\(\beta\) [56]) are essential for viability of bloodstream \(T. brucei\). Our discovery that these essential enzymes bind 4-anilinoquinazoline drugs that kill the parasite suggests that this class of small molecules is a valuable starting point for medicinal chemistry optimization [63–65] as anti-trypanosomal agents. That work is in progress.

A General Method for Discovering Targets of Kinase Inhibitor Drugs

Lapatinib kills bloodstream \(T. brucei\) with a GI\(_{50}\) in the low micromolar range (Fig. 2). To advance studies aimed at optimizing anti-trypanosome properties of lapatinib, it is important to identify the protein kinases that recognize the drug. This task is made challenging by the absence of EGFR/HER2 family of kinases in \(T. brucei\) [19,52]. Chemical proteomics [66] involving drug affinity chromatography seemed a reasonable path to discover the trypanosome targets for lapatinib. Nevertheless, we opted to develop a more versatile protocol that would not require the use of specialized matrices containing covalently attached kinase inhibitors (e.g., kinobeads) [66,67].

Since all protein kinases can bind to ATP, one could use the trinucleotide as a universal ligand for that class of enzymes (as well as other ATP-binding proteins). Therefore we used an ATP-affinity column, instead of covalently attaching the drugs directly to a matrix. After that we bound a lysate of total cellular proteins to the affinity column, we washed it stringently to disrupt non-specific protein-matrix interactions, and then eluted with drugs (e.g., lapatinib or canertinib or AEE788). Proteins eluted by the drugs were identified with mass spectrometry and bioinformatic approaches (see Materials and Methods) (see Table 1). Our approach can be used to discover cellular targets of drugs that interfere with ATP-protein interactions/metabolism. Indeed we have used the protocol to discover trypanosome protein targets of drugs other than those reported here (R. Behera and K. Mensa-Wilmot, in preparation.). A related technique was published while this manuscript was being written [68].

It is worth mentioning that the protocol devised for this study parallels the situation that a drug encounters intracellularly where ATP is present at millimolar levels. Our protocol replicates this scenario by adding a cell lysate to an ATP affinity column first, before introducing a drug to elute the target protein. ATP
Four Trypanosome Proteins are Likely to Adopt a Lapatinib-Compatible Backbone Conformation

Lapatinib is one of the most selective among 38 kinase inhibitors tested against 317 human enzymes [57]. In addition to the Tyr kinases EGFR/HER2, the drug binds with appreciable affinity to three other protein kinases, namely STK10/LOK, RIPK2, and STK2/SKL [70–72]. This high selectivity of lapatinib is partially due to a specific conformation of the kinase domain bound by the drug [32]; it is characterized by (i) a broken conserved Lys-Glu salt bridge, (ii) a displaced βC helix, and (iii) a flipped conserved Phe at the base of the “activation loop” (i.e., DFG motif) [64,73,74].

Up to 45% of kinases in the PDB, according to our estimates, can adopt a lapatinib-compatible backbone conformation (unpublished). Many structures of proteins with high sequence homology can adopt a lapatinib-compatible EGFR template (PDB 1xkk) produced structures that are free of steric conflicts or energetically strained regions (Fig. 4). These modeling data support conclusions that trypanosome lapatinib-binding proteins can adopt a lapatinib-compatible backbone as one of the equilibrium conformations for their kinase domains (Fig. 5).

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