An Ebola, Neisseria and Trypanosoma human protein interaction census reveals a conserved human protein cluster targeted by various human pathogens

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A B S T R A C T

Filovirus ebolavirus (ZE; Zaire ebolavirus, Bundibugyo ebolavirus), Neisseria meningitidis (NM), and Trypanosoma brucei (Tb) are serious infectious pathogens, spanning viruses, bacteria and protists and all may target the blood and central nervous system during their life cycle. NM and Tb are extracellular pathogens while ZE is obligatory intracellular, targeting immune privileged sites. By using interactomics and comparative evolutionary analysis we studied whether conserved human proteins are targeted by these pathogens. We examined 2797 unique pathogen-targeted human proteins. The information derived from orthology searches of experimentally validated protein–protein interactions (PPIs) resulted both in unique and shared PPIs for each pathogen. Comparing and analyzing conserved and pathogen-specific infection pathways for NM, TB and ZE, we identified human proteins predicted to be targeted in at least two of the compared host-pathogen networks. However, four proteins were common to all three host-pathogen interactomes: the elongation factor 1-alpha 1 (EEF1A1), the SWI/SNF complex subunit SMARCC2 (matrix-associated actin-dependent regulator of chromatin subfamily C), the dolichyl-diphosphooligosaccharide–protein glycosyltransferase subunit 1 (RPN1), and the tubulin beta-5 chain (TUBB). These four human proteins all are also involved in cytoskeleton and its regulation and are often addressed by various human pathogens. Specifically, we found (i) 56 human pathogenic bacteria and viruses that target these four proteins, (ii) the well researched new pandemic pathogen SARS-CoV-2 targets two of these four human proteins and (iii) nine human pathogenic fungi (yet another evolutionary distant organism group) target three of the conserved proteins by 130 high confidence interactions.

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1. Introduction

About 10 13 human cells coexist with 10 14 bacterial, fungal, and protozoan cells, i.e., thousands of microbial species. This means that we live in “close intimacy” with a wide variety of microbes. Some of them are capable of causing us illness or death once the “eco-system” is broken and an immunocompromised situation arises, as is the case for Neisseria meningitidis [1]. For the aims of this manuscript we will name it as an opportunistic pathogen. On the other hand, dedicated pathogens do not require the host to be immunocompromised or injured before invading it. Their specialized mechanisms for crossing cellular and biochemical barriers and for eliciting specific responses from the host organism contribute to the survival and multiplication of the pathogen, as is the case of Ebola virus and Trypanosoma sp. The host responses may even contribute to pathogen survival and multiplication, as happens for Ebola virus and Trypanosoma sp. [1–5].

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Furthermore, (i) viruses, including Ebola virus, only replicate inside cells, thus being intracellular pathogens; (ii) the Gram-negative bacterium \(N. meningitidis\) is an asymptomatic colonizer (harmless commensal) of the human nasopharynx, mostly using propionic acid available outside of cells in that environment. However, to cause disease it must traverse the blood-cerebrospinal fluid (B-CSF) barrier. This has lead researchers to postulate transcellular passage (and, implicitly, cell entry); (iii) the parasitic protist \(T. brucei\) is not intracellular in either of its two hosts, tse-tse flies (insect stage) or man/cattle (bloodstream stage); however, the constantly changing specific factor variant surface glycoprotein (VSGs) of \(T. brucei\) constitutes a complicating issue. How trypanosomes cross the brain parenchyma and persist behind an intact blood–brain barrier (BBB), how their growth in the brain is controlled and how the unicellular trypanosomes exit the brain and cause a relapse are still questions to be answered. Additional details can be found in Supplementary file 1.

Cohabitation between human populations at risk and their animals - as pathogen reservoirs - is common, and their everyday contact may even permit disease transmission [6]. Thus, ecological and evolutionary hints, together with disease dynamics, are fundamental for understanding and analysing the temporal, organizational and spatial interactions, even within a single pathogen. That is, hours to months, cellular to ecosystem levels, and local to pandemic spread of diseases [6]. Additionally, some pathogens circulate between individuals of a single species, while others flow among multiple hosts, need insect vectors, or subsist in reservoirs. Hence, we compare in this first part of our study Filovirus ebolavirus (ZE for Zaire ebolavirus, Bundibugyo ebolavirus), \(N. meningitidis\) (NM for \(N. meningitidis\)) and \(T. brucei\) - expressing a miriad of host-pathogen interactions.

It is fundamental and may be attractive to elucidate defense strategies common among hosts when confronted with these three diverse pathogen types - whether opportunistic or professional - expressing a miriad of host-pathogen interactions. Interactions and strategies may differ, and yet investigating those commonly utilized would help further define how the immune response reacts to pathogens. However, we realized during the course of this study that the tools we have at our disposal are only \textit{in silico} and to uncover immune reactions to pathogens would require a large-scale study with many organisms and many experiments testing human immune responses. Instead, to validate our findings from these three very different organisms, we can with our \textit{in silico} approach only systematically determine how far always the same human proteins are targeted by a wide variety of pathogens while any functional conclusions need experimental data. For both parts of our study we used a host-pathogen protein–protein interactions (HP-PPIs) framework [7]. On a large scale, HP-PPIs systematically provide a global view to understand the basis of infectious diseases while at small-scale it is possible to identify conserved interactions in different host-pathogen pairs. PPIs conservation reflects the different levels of interactions paving the route to infection and host defense response. The aim, design and setting of the study were to seek direct information on the proteins and protein interactions in pathogens targeting human proteins from different kingdoms to reveal potential conserved host interactions as well as specific or only partly shared protein interactions.

However, only a systematic approach is here powerful, hence we rechecked and could then confirm the most central four human proteins commonly targeted in all three pathogens by a systematic census over human pathogenic organisms, comparing bacteria, viridae and fungi. The available host-pathogen protein–protein interaction databases support conserved targeting of these four human proteins by various human pathogens.

### 2. Methods

The flow diagram (Fig. 1) illustrates all the pipeline steps.

#### 2.1. Proteomes and amino acid sequences

The reference proteomes of human (reference proteome accession: UP000005640) and \(Bundibugyo ebolavirus\) (reference proteome accession: UP000143891) were retrieved from Uniprot database [8]. The reference proteomes of \(Trypanosoma brucei brucei\) TREU927 (genome assembly ASM244v1, RefSeq assembly accession: GCF_000002445.1) and \(Neisseria meningitidis\) MC58 (genome assembly ASM880v1, RefSeq assembly accession: GCF_000008805.1) were retrieved from RefSeq database [9].

#### 2.2. Inferring protein–protein interactions based on interologs

The experimentally determined host-pathogen PPIs from PHISTO database [10] and PPIs present in the DIP (Database of Interacting Proteins) database [11] were used as a template to reconstruct interologs based PPIs networks [12] of human-ZE, human-Tb and human-NM. The stand-alone InParanoid program version 4.1 was used to assign proteins from the three pathogens to orthology groups. It uses the reciprocal best blast hit strategy to identify the clusters of orthologs and paralogs [13]. Only the seed ortholog pair whose Bootstrap value was 100% was considered to increase prediction confidence.

#### 2.3. Pruning PPIs with domain-domain interactions

The amino acid sequences of identified non-redundant DIP-based interologs of host-pathogen PPIs were extracted and domains were assigned to the sequences using Pfam version 29.0 [14]. The information of interacting protein domains was prepared using the three databases Domine [15], DIMA 3.0 [16] and IDDI database [17]. They consider the information of the Pfam families suggested to interact based on structural information present in the protein complexes available in Protein Data Bank (PDB) [18]. This list was used to parse the DIP-based interologs of PPIs using customized Perl scripts, and only those interolog [12,19] interactions consisting of at least one true domain-domain interaction pair as listed in the chosen databases were retained.

#### 2.4. Host-pathogen PPI networks

The host-pathogen PPIs predicted from both the PHISTO and DIP-based interactions were merged and redundancies were removed. Cytoscape version 2.8.1 [20] was used to visualize all three host-pathogen PPI networks. The network hubs were determined using the cyto-Hubba plugin [21]. The networks were further curated by extensive manual effort, and the connectors of proteins related to the human immune system were retrieved to generate three immune subnetworks. Immune subnetworks were further analysed with Cytoscape to identify the central host proteins targeted by all three selected pathogens.

#### 2.5. Gene-Ontology (GO) annotation and pathway Over-Representation analysis (ORA)

The protein sequences of pathogen interacting host proteins were extracted from all three established host-pathogen PPIs and resulting proteins were functionally annotated by the BLAST2GO software suite version 2.4.1 [22]. Fisher’s exact test was used to identify significantly over-represented GO terms. The pathway over-representation analysis of conserved human proteins that
were targeted by at least two pathogens was performed using the pathway analysis tool at InnateDB database [23]. The p-value was calculated using hypergeometric tests, and Benjamini-Hochberg adjustment was used for multiple test correction.
2.6. Subcellular localization

The subcellular localization of immune-related human proteins was determined with extended version of KnowPredsite [24] available at UniLoc server (bioapp.iis.sinica.edu.tw/UniLoc/), a knowledge-based classifier for protein subcellular localization [25]. The prediction server KnowPredsite II is capable of predicting single and multiple localization sites for both the eukaryotic and prokaryotic proteins.

2.7. Analysis of conservation of host-pathogen protein interactions and protein targets

All host-pathogen protein–protein pairs and interactions from the HPIDB database [26] were examined and conservation of identified interactions as well as conserved targeting of proteins by all three pathogens were accessed. We determined conservation of interactions using the interolog approach [19]. We further assessed the conservation of targeting by further organisms regarding the identified common host proteins. For this validation we collected the interaction data from the PHISTO database [10], human-SARS-CoV-2 interactomes [27,28], and FungiWeb database (https://fungiweb.bioapps.biozentrum.uni-wuerzburg.de/php/fungiweb.php) and analysed the targeting of identified conserved host proteins comparing different pathogens.

3. Results

3.1. Reconstruction of host-pathogen interaction networks

Although primary interactions between host cells and pathogens are usually diverse, pathogens use several mechanisms to signal their presence or to invade host cells. As these host-pathogen PPIs mainly occur due to biophysical properties of the involved protein molecules, they do not completely disrupt host protein networks. They can, however, effectively modulate host signaling and interactome [29,30]. Of note, critical proteins from the host interactome [31] which enable the pathogen to switch host systems [32] are preferentially targeted. We reconstructed and analyzed HP-PPI (host-pathogen protein–protein interaction) networks comparing the virus Zaire ebolavirus, (Bundibugyo ebolavirus, ZE), the prokaryotic bacterium Neisseria meningitidis (NM) and the eukaryotic protist Trypanosoma brucei gambiense and rhodesiense, (Tb). We mapped physical PPIs between the human host and each pathogen, using orthology relations and host-pathogen interaction databases. Briefly, to reconstruct these HP-PPI networks, we first collected each pathogen and human genome and their corresponding protein-coding genes. Next, we created the respective networks using PHISTO and DIP database as templates of experimentally verified PPIs (see material and methods section). Due to their structural features, exposed protein domains have higher probabilities to be involved in PPIs. These protein domains are often conserved [33] and used by many pathogens to invade and disseminate in the host [34,35]. Therefore, we used a stringent interaction mapping pipeline that refines contact predictions by considering known domain-domain interactions. This approach produces high-quality interactome data with low false-discovery rates [36–39]. The list of all obtained interactions is given in Supplementary file 2.

To further characterize protein functions in these host-pathogen PPI networks we used Gene Ontology (GO) annotation. Fig. 2 displays the number of genes involved in the interactome for each pathogen and its counterpart in humans (panels a-d). Although each HP-PPI network express unique characteristics, the overall profile of gene type involved in the interaction is similar for all three classes of organism analyzed. That is, most host proteins involved in the interaction are expressed either in the plasma membrane, or relate to DNA or RNA functions, or are mitochondrial proteins. Furthermore, with increasing complexity (from viruses to lower eukaryotes) of the invading pathogen, an increasing number of proteins seem to be involved in the host-pathogen interaction for all compartments or functions; of note, most pathogen proteins participating in the interaction are not well characterized in their function. We hence focused in the second part of our investigation more on the conservation of the targeting of the well characterized human proteins by various pathogens.

Fig. 2g and 2h compare the number and percentage of human genes related to each category participating in the interactome for each disease. Fig. 2i and 2j illustrate the number and percentage of pathogen genes related to each category involved in the interactome for NM and Tb. The results indicate that the overall profile of protein type contributing to the interaction is similar and does not depend on which organism class the pathogen belongs, thus suggesting a high conservation involved in these host-pathogen interactions.

3.2. Unique and shared human proteins in host-pathogen networks

Pathogens tend to interact with host network hubs hijacking them for their own profit [40]. However, the host network should remain “robust”, to prevent catastrophic failure of host cell metabolism. The reverse would hinder pathogen survival and proliferation and kill the host [41]. We thus analyzed proteins participating in the interactome and represented in the HP-PPI networks. Supplementary file 3 lists pathogen targeted human proteins by all three pathogens. Supplementary file 3 lists pathogen targeted human proteins in reconstructed HP-PPI networks. A total of 2797 unique human proteins were targeted by the three analyzed pathogens. Eukaryotic cells herein represented by trypanosomes were found to achieve the highest numbers of specific host interactions (1414); this was followed by Neisseria (753) and Ebola (57), respectively. To dissect paths involved in each host-pathogen interaction, we further performed pathway analysis and classification, collected by approaches such as Netpath [42], BioCarta [43], KEGG [44] and Reactome [45] database classification.

Analysis of the human-Tb HP-PPI network using the Netpath database identifies EGFR, thymic stromal lymphopoietin (TSLP) and androgen receptor pathways, while data collected from Reactome point towards host defense mechanisms, immune signalling, cell senescence, and response to stress as top categories involved. According to the BioCarta database, further signalling relies mainly on thrombospondin-1 (Tsp-1) induced apoptosis, extracellular-signal-regulated kinases/FYN proto-oncogene, ssrc family tyrosine kinase/Tyrosine kinase (Erb/Fyn/Tyr) activation and EGF signalling. These results are further supported by KEGG which identifies NOD-like receptor signalling, antigen processing, and trypanosomal defense pathways, as central for the HP-PPI network. Finally, InnateDB classifies innate responses against trypanosomes as relying on IL-2 and IL-12 as well as osteopontin and atypical NF-kB mediated processes (Supplementary file 4).

Despite targeting common proteins, host defense details vary for each disease. For example, interacting host proteins shared between NM and ZE include those associated with mRNA stability, gene expression control, and RNA processing and splicing (reactome classification). Remarkably, Netpath reveals again androgen receptor pathways as significant, and TNFalpha response pathways as involved in host response against these two pathogens.

According to the reactome database, ZE and Tb, despite differing greatly in complexity and size, trigger shared specific human defense systems centering on eukaryotic translation, elongation,
life cycle, and infection, involving ribosomes and PID-Bard1 signalling, modulated by Rb and p53 proteins according to KEGG. Additionally, Netpath shows a shared TSLP pathway response and again the TNFalpha response.

As a conclusion from this triple comparison, our data suggest that commonly targeted host proteins include host defense and differentiation/maturation pathways. Additionally, the two bigger HP-PPI networks (NM, Tb) also involve reprogramming of host-cell senescence and apoptosis; the two smaller pathogens (NM, ZE) influence mRNA; and Tb and ZE impact protein translation and specific host defense pathways.

3.3. Characterization of pathogen proteins and their role in human-pathogen interaction

We further reviewed disease characteristics for ZE, NM and Tb. Detailed information is listed and explained in Supplementary file 5 (Comparison Table) and Supplementary file 1 (Text file including details for each pathogen and disease), including general responses occurring during the infection processes of all three pathogens. A thorough assessment of the summarized data points provides functional similarities and differences occurring during infection processes. The collected biological information on immune-
related and infection-associated proteins was used for extensive individual curations of the reconstructed preliminary HP-PPI networks. This literature-driven analysis was crucial to address the next phase of our investigation directed towards the identification of infection-related HP-PPIs, based on these curated subnetworks and comparisons between host and pathogen interactions for each organism.

Identification of specific infection-related HP-PPIs and their association with data presented in Supplementary file Figure 1 to 3; Supplementary data 4 till 12 (comparing and describing disease characteristics for each pathogen) allowed the description of useful interaction examples.

For example, the Ebola virus glycoprotein ZE-GP, enables viral entry into target cells [46]. It also antagonizes host cell protein tetherin (bone marrow stromal antigen 2, BST2, also known as tetherin, see Table 1) antiviral activity. This plasma membrane molecule usually restricts virus release from infected cells. It is not clear how ZE-GP antagonizes tetherin. However, this result emphasizes the interaction existing between both molecules and highlights that understanding how Ebola virus offsets innate immune system antiviral effectors might help to define novel antiviral intervention targets. BST2 plays a key role in cellular antiviral response against several enveloped virus including HIV-1, thus preventing subsequent infections [47,48]. BST2 has recently

Fig. 3. Unique and common targeted human proteins by three pathogens: Neisseria meningitidis, Trypanosoma brucei and Zaire ebolavirus. See Supplementary file 3 for annotation of involved proteins.
| Zealomonas sp. NAME | Zealomonas sp. NAME | Zealomonas sp. NAME |
|---------------------|---------------------|---------------------|
| ZeaMaxX | ZeaMaxX | ZeaMaxX |
| ZeaMaxX | ZeaMaxX | ZeaMaxX |
| ZeaMaxX | ZeaMaxX | ZeaMaxX |

### Table 1

Host defense associated human proteins involved in physical interaction with Zea, TB and NM proteins.

| ZeaMaxX | ZeaMaxX | ZeaMaxX |
|---------------------|---------------------|---------------------|
| ZeaMaxX | ZeaMaxX | ZeaMaxX |
| ZeaMaxX | ZeaMaxX | ZeaMaxX |
| ZeaMaxX | ZeaMaxX | ZeaMaxX |

Color code: Interferon associated proteins, Blue: Antigen, antibody and HLA associated proteins, Green: Cytokine and chemokine associated proteins, Red: Proteasome associated proteins, Light brown: Transcription factors associated proteins, Red: Cell recognition associated proteins. Note that EBOV-human immune system interaction concentrates in two proteins, one at the nuclear level (proliferation) and one that can be found in the cytoplasm, Golgi apparatus and plasma membrane. On the other hand, human immune related proteins involved in the interaction with T. brevi and N. meningitidis overlap in high proportion. In the table the following codes are used: U for unique, and C for conserved interactions; the numbers correspond to the number of pathogenic proteins in corresponding category interacting with them. Information about the function of these proteins can be found in the Supplementary file 6. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
caught attention as deletion of BST2 cytoplasmic and transmembrane N-terminal domains results in SARS-CoV, SARS-CoV-2, and influenza virus production suppression in a VERO cell line [49]. Hence, this host protein is important for Ebola virus replication and potentially for several corona viruses. BST2 has been implicated in SARS-CoV-2 cellular restriction [50] and SARS-CoV-2 spike protein generally downregulates tetherin to enhance viral spread [51].

A second example related to NM follows. Heptose 1,7 bisphosphate (HBP) actively secreted or released after NM lysis seems to be delivered to the cytoplasm through the endocytic route. HBP is a metabolic intermediate in lipopolysaccharide (LPS) biosynthesis and is highly conserved in Gram-negative bacteria. HBP presence in the cytoplasm is sensed by tumor necrosis factor receptor-associated factor (TRAF)-interacting protein with a forkhead-associated domain (TIFA). During infection, signaling series of events occur including TRAF2/6 [TNF receptor-associated factor 6 (Table 1) oligomerization and ubiquitination. After recruitment and inclusion of additional host factors this molecular signaling leads to the activation of nuclear factor (NF)-kB and proinflammatory gene expression [52,53].

A final example (Supplementary files 1 and 5) highlights that in trypanosomiasis, several genes and pathways associate with the latent phenotype, suggesting an increased activation of both T- and B-cells in patients with latent infection [54]. The data suggest increased lymphocyte activation in HAT patients, relative to latent cases. Additional information also predict activation of lymphocytes (p < 0.0001), activation of T cells (p = 0.0004), co-stimulation of T lymphocytes (p = 0.0013), and Ca²⁺ flux (p = 0.0004) as an indication of lymphocyte activation, as significantly different between active and latent cases of trypanosomiasis. Using transcriptome data to conduct an exome-wide single nucleotide polymorphism (SNP) association has suggested that SNP in human major histocompatibility locus associates with disease severity, thus suggesting that T cell activation is a determining factor in disease outcome [54].

The overlapping curated HP-PPI subnetworks of interacting proteins belonging to pathogen and host were further used to understand the role of described host defense PPIs. Subnetwork comparison suggested the existence of common human mechanisms to fight against these three pathogens as depicted in Fig. 4-a-c and summarized in Table 1 (detailed interactions are listed in Supplementary file 6). Fig. 4 illustrates how redundancy of host nodes in different HP-PPI networks are fundamental for survival during host-pathogen encounters [55], either to impair or to promote host protection, relying on innate and cellular defense mechanisms [56]. Moreover, a direct mapping of functional processes (Supplementary files 1 and 5), can be built from processes and gene

Fig. 4. An interaction map of the human host with three pathogens comparing protein–protein interactions (PPIs). The map is based on interologs. The color of the nodes indicates the different organisms: (a) Neisseria meningitidis = red; (b) Trypanosoma brucei = blue; (c) Zaire ebolavirus = yellow, and green = human proteins. Human nodes are shaded in the color clouds and arranged according to their subcellular localization. The border thickness around green human nodes indicates the confidence of localization in different cellular compartments. Membrane proteins of N. meningitidis are depicted by underlined and italics text while the N. meningitidis membrane proteins with multiple localization is shown in italics. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)
Fig. 4 (continued)
ontology associated with these protein networks (listed with uniport identifiers and annotations both for human host and pathogen in Supplementary file 6). Additionally, an analysis of strongly connected (protein hubs) or weakly connected proteins can be found in Supplementary file 7. In this way a systematic overview of proteins, annotations and processes is given, appropriately fitting disease characteristics and considering the identified human host and pathogen protein interactions.

Host-pathogen interaction details and individual protein function including host defense-related signalling pathways and compartments are given in Table 1. This table categorizes unique (U) and conserved (C) host proteins involved in the interaction with pathogens. Specific host protection mechanisms against all three pathogens include interferon associated and induced proteins, the HLA system, cytokines and chemokines and proteasome associated proteins. Regulation of this host defense network is achieved by transcription factors (yellow lettering) via cell recognition associated proteins (red lettering). Interestingly, the ZE interaction molecules interact strongly with two human proteins, one is located in the nucleus and related to proliferation) and another one has multiple locations including cytoplasm, Golgi apparatus, and plasma membrane: lipid raft associated protein CD antigen CD317, or stromal bone marrow antigen-2, respectively (see Fig. 4).

3.4. Central interactome proteins shared by all pathogens and humans, implications

Our HP-PPI results illustrate that only four host proteins targeted all three HP-PPI networks: the elongation factor 1-alpha 1 (EEF1A1), the SWI/SNF complex subunit SMARCC2, the dolichyl-diphosphooligosaccharide--protein glycosyltransferase subunit 1 (RPN1), and the tubulin beta-5 chain (TUBB). The identified host-pathogen interactions as compared in Fig. 4a-c, suitable for each specific pathogen disease characteristic (supplementary files 1 and 5), share a common denominator: top four hub proteins are exploited by pathogens to gain entry to the host and survive inside it. As illustrated in Fig. 5, (colour codes differentiating all three pathogens), these four central proteins are strongly and well connected in the human interactome (Supplementary Fig. 1). Nor-
mally, hub proteins are well conserved [19,57] and hence represent good pathogen targets [29].

Based on these results that suggest a central role of these host hub proteins, we postulate that they may represent potential targets for the design of infection disruption tools and that their manipulation might stop all three infections. For example, EEF1A1 relates to host cell functional changes, so that infection can better exploit cell protein synthesis. SMARCC2 and the TUBB may target cytoskeleton hubs related to infection; their modulation may block all three infections. RPN1 might provide an important target involving glycosyl and lipid metabolism, which all of the three infective agents use.

Table 2 lists these conserved four human proteins as well as their interactions with pathogen proteins, inferred by homology, predicted, putative, or demonstrated by transcription. Involved functions and processes of these proteins are also described in Table 2. Accumulating experimental evidence suggests that these pathogen proteins share these central interactions with the host and exploit this for pathogen entry. In particular for Ebola virus, the interactions have been demonstrated experimentally by high-affinity purification coupled to label-free mass-spectrometry-based approach for VP24 and all four human proteins [58,59]. Moreover, further evidence is available for Trypanosoma and even in selected cases for Neisseria (Table 2). Altogether, these results suggest that a common, shared protein interaction route could be implicated in pathogen entry.

3.5. Large-scale host-pathogen data validate the clade of four host proteins to be well conserved among different organisms in host-pathogen interactions

To validate this unexpected finding further, we analyzed conservation of these four central interactions in large-scale host pathogen databases over animals, fungi, bacteria and viruses and checked participation of all four host proteins in different cellular pathways and networks.

3.5.1. Experimental data confirming the pathogen protein interaction with four conserved human proteins

The conservation of the identified host-pathogen interactions can not be accurately accessed because of the lack of experimentally validated interspecies interactions between pathogens and host species other than human. The largest collection of interspecies host-pathogen interaction database (HPIDB) contains 94.6% exclusively interactions between pathogens and humans [26] (Supplementary file 8a). The second most common host species in HPIDB is mouse which consists of only 1.1% of interspecies interactions. The rest of the 64 host species contributes 4.2% of total interactions. This bias towards human data do not allow us to validate interaction conservation comparing multiple host-pathogen interaction pairs. Therefore, we assessed the identified interactions with four central proteins (EEF1A1, RPN1, SMARCC2, TUBB) independently (Table 3). The interaction between all the four central proteins and ZE protein VP24 has been validated by tandem affinity purification experiments [58]. In total, two host-pathogen interactions involving EEF1A1 and NM, one interaction involving RPN1 and NM, two interactions involving SMARCC2 and NM, one interaction involving TUBB and NM were transferred from highly conserved experimentally verified host-pathogen interactions. Similarly, two interaction involving EEF1A1 and Tb and 1 interaction involving TUBB and Tb were transferred from highly conserved experimentally verified host-pathogen interactions. We identified unique interaction between EEF1A1 and Tb, three interactions between RPN1 and Tb, and one interaction between SMARCC2 and Tb by interaspecies interaction transfer and domain-domain interactions. All these mentioned interactions are listed in (Table 3).

3.5.2. Many pathogenic bacteria and viruses target these four conserved human proteins

To understand the importance of these four central proteins, we studied further the conservation of its targeting by different pathogens. We used the interaction data from PHISTO database which comprise human experimental interactions with 97 bacteria, 6 fungi, 11 protozoa and 474 viruses [10] and we examined whether and to which extent these pathogens target the four central proteins identified by our analysis. This assessment revealed altogether 56 unique pathogen interactions from bacteria and viruses that target these four central proteins to establish the infection (Supplementary file 8b). For instance, EEF1A1 was targeted by 36 unique pathogen species/strains creating 60 host-pathogen interactions (Fig. 6).

This clearly indicates that these four central proteins are among the preferential targets of pathogens.

3.5.3. SARS-CoV-2 targets also two proteins of this conserved protein clade

To examine the importance of our four human proteins further we also checked whether these proteins are targeted by the novel, intensely investigated global human pathogen SARS-CoV-2 by analysing published, experimentally confirmed human-SARS-CoV-2 interactomes [27,28]. We found experimental evidence that out of these four protein targets, EEF1A1 and RPN1 are also targeted by SARS-CoV-2 (Supplementary Fig. 2).

In a recent work, EEF1A1 was shown as a potential drug target against the B.1.1.7 variant of SARS-CoV-2 [60]. EEF1A1 can be efficiently targeted by plitidepsin [61]. On the pathway level EEF1A1 and RPN1 both also involves in metabolism of proteins and translation (Supplementary file 8c) and hijacking the host translational machinery is one the survival strategy of SARS-CoV-2 [62].

3.5.4. Various pathogenic fungi target these four conserved human proteins

We further looked at the FungiWeb database to identify the interaction between proteins of nine fungal pathogens (Aspergillus fumigatus, Candida glabrata, Candida albicans, Coccidioides immitis, Cryptococcus neoformans, Debaryomyces hansenii, Lodderomyces elongisporus, Paracoccidioides brasiliensis, Rhizopus delemar) with central host targets. Here we identified 130 high confidence interactions with these fungal pathogens and three of the four conserved human proteins: EEF1A1, RPN1 and TUBB (Supplementary file 8d). EEF1A1 was identified as a highly targeted protein as all the nine fungal pathogen targets this protein and all together creates 78 interactions (Supplementary Fig. 3). Pathway analysis of these four proteins (Supplementary file 8c) reveals that all four are involved in cytokskeletal signaling and regulate by this also other central processes such as translation and host cell entry (EEF1A1), protein processing, proteasome and translation (RPN1), chromatin remodeling (SMARCC2), cell cycle and centrosome maturation (TUBB).

Taken together, these three interaction surveys confirm that the four proteins are often targeted by various human pathogens, even if they are far evolutionary apart and from different kingdoms (see discussion).

4. Discussion

Pathogens infect and replicate and grow inside the human host. Targeting host proteins improves survival in the host and is selected in evolution. We wanted to better understand specific interactions but so far the available methods to do so are not optimal. To address this, we developed a large-scale analysis of human-pathogen interactions (HPI) and took advantage of a merge of three different methods 3.5.1. Experimental data confirming the pathogen protein interaction with four conserved human proteins

The conservation of the identified host-pathogen interactions can not be accurately accessed because of the lack of experimentally validated interspecies interactions between pathogens and host species other than human. The largest collection of interspecies host-pathogen interaction database (HPIDB) contains 94.6% exclusively interactions between pathogens and humans [26] (Supplementary file 8a). The second most common host species in HPIDB is mouse which consists of only 1.1% of interspecies interactions. The rest of the 64 host species contributes 4.2% of total interactions. This bias towards human data do not allow us to validate interaction conservation comparing multiple host-pathogen interaction pairs. Therefore, we assessed the identified interactions with four central proteins (EEF1A1, RPN1, SMARCC2, TUBB) independently (Table 3). The interaction between all the four central proteins and ZE protein VP24 has been validated by tandem affinity purification experiments [58]. In total, two host-pathogen interactions involving EEF1A1 and NM, one interaction involving RPN1 and NM, two interactions involving SMARCC2 and NM, one interaction involving TUBB and NM were transferred from highly conserved experimentally verified host-pathogen interactions. Similarly, two interaction involving EEF1A1 and Tb and 1 interaction involving TUBB and Tb were transferred from highly conserved experimentally verified host-pathogen interactions. We identified unique interaction between EEF1A1 and Tb, three interactions between RPN1 and Tb, and one interaction between SMARCC2 and Tb by interaspecies interaction transfer and domain-domain interactions. All these mentioned interactions are listed in (Table 3).

3.5.2. Many pathogenic bacteria and viruses target these four conserved human proteins

To understand the importance of these four central proteins, we studied further the conservation of its targeting by different pathogens. We used the interaction data from PHISTO database which comprise human experimental interactions with 97 bacteria, 6 fungi, 11 protozoa and 474 viruses [10] and we examined whether and to which extent these pathogens target the four central proteins identified by our analysis. This assessment revealed altogether 56 unique pathogen interactions from bacteria and viruses that target these four central proteins to establish the infection (Supplementary file 8b). For instance, EEF1A1 was targeted by 36 unique pathogen species/strains creating 60 host-pathogen interactions (Fig. 6).

This clearly indicates that these four central proteins are among the preferential targets of pathogens.

3.5.3. SARS-CoV-2 targets also two proteins of this conserved protein clade

To examine the importance of our four human proteins further we also checked whether these proteins are targeted by the novel, intensely investigated global human pathogen SARS-CoV-2 by analysing published, experimentally confirmed human-SARS-CoV-2 interactomes [27,28]. We found experimental evidence that out of these four protein targets, EEF1A1 and RPN1 are also targeted by SARS-CoV-2 (Supplementary Fig. 2).

In a recent work, EEF1A1 was shown as a potential drug target against the B.1.1.7 variant of SARS-CoV-2 [60]. EEF1A1 can be efficiently targeted by plitidepsin [61]. On the pathway level EEF1A1 and RPN1 both also involves in metabolism of proteins and translation (Supplementary file 8c) and hijacking the host translational machinery is one the survival strategy of SARS-CoV-2 [62].

3.5.4. Various pathogenic fungi target these four conserved human proteins

We further looked at the FungiWeb database to identify the interaction between proteins of nine fungal pathogens (Aspergillus fumigatus, Candida glabrata, Candida albicans, Coccidioides immitis, Cryptococcus neoformans, Debaryomyces hansenii, Lodderomyces elongisporus, Paracoccidioides brasiliensis, Rhizopus delemar) with central host targets. Here we identified 130 high confidence interactions with these fungal pathogens and three of the four conserved human proteins: EEF1A1, RPN1 and TUBB (Supplementary file 8d). EEF1A1 was identified as a highly targeted protein as all the nine fungal pathogen targets this protein and all together creates 78 interactions (Supplementary Fig. 3). Pathway analysis of these four proteins (Supplementary file 8c) reveals that all four are involved in cytokskeletal signaling and regulate by this also other central processes such as translation and host cell entry (EEF1A1), protein processing, proteasome and translation (RPN1), chromatin remodeling (SMARCC2), cell cycle and centrosome maturation (TUBB).

Taken together, these three interaction surveys confirm that the four proteins are often targeted by various human pathogens, even if they are far evolutionary apart and from different kingdoms (see discussion).

4. Discussion

Pathogens infect and replicate and grow inside the human host. Targeting host proteins improves survival in the host and is selected in evolution. We wanted to better understand specific interactions but so far the available methods to do so are not optimal. To address this, we developed a large-scale analysis of human-pathogen interactions (HPI) and took advantage of a merge of three different methods
| Human protein | UniProt ID | Human protein function | Pathogen | Protein from pathogen | Uniprot ID (evidence level)* | Pathogen protein function | Reported experimental validation | Reference |
|---------------|------------|------------------------|----------|-----------------------|----------------------------|---------------------------|-------------------------------|-----------|
| Dolichyl-diphosphooligosaccharide–protein glycosyltransferase subunit 1 | P04843 | Post-translational protein modification | *Trypanosoma brucei* | DNA polymerase kappa | Q384G6 (predicted) | Error-prone translesion synthesis | N/A | [80]** |
| | | | *Trypanosoma brucei* | Endoplasmic reticulum oxidoreductin | Q571U3 (predicted) | Protein folding | N/A | |
| | | | *Trypanosoma brucei* | Protein transport protein Sec13 | Q38B44 (inferred from homology) | Protein transport from endoplasmic reticulum DNA replication, error-prone translesion synthesis | N/A | |
| | | | *Neisseria meningitidis* | DNA polymerase IV | Q9JYS8 (inferred from homology) | Host-virus interaction, influencing host immune response | N/A | [59] |
| | | | *Ebola virus* | Membrane-associated protein VP24 | Q9J322 | affinity chromatography technology | N/A | |
| SWI/SNF complex subunit SMARC2 | Q8TAQ2 | Chromatin remodeling | *Trypanosoma brucei* | Aldehyde dehydrogenase | Q38AY7 (putative) | Host-virus interaction, influencing host immune response | N/A | |
| | | | *Neisseria meningitidis* | Succinate-semialdehyde dehydrogenase | Q9JYP4 (inferred from homology) | Catalysis aldehyde oxidation | N/A | |
| | | | *Neisseria meningitidis* | Ferrochelatase | Q9K097 (inferred from homology) | Cell grown in presence of Mn | N/A | |
| | | | *Ebola virus* | Membrane-associated protein VP24 | Q9J322 | Porphyrin biosynthesis | N/A | |
| | | | *Neisseria meningitidis* | Biosynthetic arginine decarboxylase | Q26802 (transcription) | Host-virus interaction, influencing host immune response survival of *Trypanosoma* | N/A | [78]** |
| | | | | | Q9K0U3 (inferred from homology) | Agmatine biosynthesis | N/A | [77,85]** |
| Tubulin beta-5 chain | P07437 | The major component of microtubules (cytoskeleton formation) | *Trypanosoma brucei* | Mitogen-activated protein kinase | Q382L7 (predicted) | Host-virus interaction, influencing host immune response | N/A | |
| | | | *Neisseria meningitidis* | Biosynthetic arginine decarboxylase | Q566B8 (inferred from homology) | Transcription regulation | N/A | |
| | | | | | | Regulating mitochondrial DNA replication | N/A | |
| | | | *Ebola virus* | Membrane-associated protein VP24 | Q26802 (transcription) | Survival of *Trypanosoma* | N/A | [78]** |
| | | | | | Q566B8 (inferred from homology) | Nucleic acid biosynthesis | N/A | |
| Elongation factor 1-alpha 1 | P68104 | Transcription regulation | *Trypanosoma brucei* | Elongation factor 1 gamma | Q382L7 (predicted) | Host-virus interaction, influencing host immune response | N/A | |
| | | | *Trypanosoma brucei* | ATP-dependent Clp protease subunit, heat shock protein 100 (HSP100) | Q566B8 (inferred from homology) | Transcription regulation | N/A | |
| | | | *Trypanosoma brucei* | Mitogen-activated protein kinase | Q26802 (transcription) | N/A | |
| | | | *Neisseria meningitidis* | Cytidylate kinase | P5706S (inferred from homology) | N/A | |
| | | | *Neisseria meningitidis* | Uncharacterized protein | Q9JYB3 | N/A | |
| | | | *Ebola virus* | Membrane-associated protein VP24 | Q9J322 | Host-virus interaction, influencing host immune response | N/A | [58] |

N/A – not available. *Uniprot accessed on July 28, 2020. **potential indirect evidence.
adaptations and common pathways shared by three dangerous human pathogens able to infect septically human beings. We compared ZE, NM and Tb and specific but also shared pathways and next test, which proteins of the host these three pathogens target. We hypothesized that this common ability of all three pathogens to be blood borne would allow to reveal ready interpretable shared host pathways attacked and found indeed four human host proteins shared as common targets by all three pathogens.

This result was then extensively counter-checked by a battery of organisms combining various data from bacterial and viral host-pathogen interaction databases, latest results from SARS-CoV-2 research as well as databases on human proteins targeted by fungi. All these data confirm: Yes, the four human proteins are well conserved and used for direct targeting by many human pathogens and we can identify the human pathways they are involved in, suggesting a central role in the infection process for the human host often suppressed by the pathogen.

4.1. A common defense network revealed from the three blood borne pathogens

4.1.1. ZE interacts with the complement cascade

Cholesterol-enriched lipid raft (plasma membrane) microdomains are a main gateway used by this virus [63,64], ZE enhances its infectivity by stimulating antibody production and virus-antibody complexes that bind to monocytes and macrophage Fc-receptors [65] and activates the complement cascade to facilitate virus entry into cells such as primate kidney cells [66]. The complex constituted by virus, antibodies and C1, then binds C1q ligands at the immune cell surface, promoting either virus binding to ZE-specific receptors or endocytosis [66].

Once internalized, a cytopathogenic effect of ZE causes annihilation of infected cells. Monocytes/macrophages and dendritic cells (DCs) are ZE early replication sites and disseminate the virus from spleen and lymph nodes to other tissues such as fibroblasts, hepatocytes, adrenal cells and epithelial cells [64]. Two proteins are involved in the ZE-human HP-PPI network as herein described: PCNA and the bone marrow stromal antigen 1 (Strain 17).

4.1.2. NM and Tb activate common pathways in the host

NM and Tb both have a blood infection stage, can invade the brain and cause meningitis. Their respective HP-PPI networks targeting human defense mechanisms overlap a lot (see Table 1), targeting antigen presentation, cytokine and chemokine secretion and antibody production. E.g. interleukin-2 receptor subunit beta, TNF receptor-associated factor 6 and interleukin-18-binding protein trigger pathogen-specific sites of thrombo-inflammation (Table 1). Extracellular matrix protein LTBP4, the latent-transforming growth factor-beta (TGF-β)-binding-protein-4, participates only in the human-NM HP-PPI network [71]. In Tb, changing glycoprotein coat or Variant Surface Glycoprotein (VSG) of bloodstream parasites is crucial for evading host antibody activity, and the complement cascade [3,72]. Similarly in NM, high-frequency antigenic variation of other surface antigens, like lipooligosaccharide (LOS), opacity-

| Host | Pathogen | Validation | Reference |
|------|----------|------------|-----------|
| EEF1A1 | VP24 | Experimentally validated | [58] |
| EEF1A1 | NMB1663 | Conserved in EEF1A1(Human)-FTT,1214c(Francisella tularensis) | [86] |
| EEF1A1 | Cmk | Conserved in EEF1A1(Human)-RCE[Bacillus anthracis] | [56] |
| EEF1A1 | TB11.01.4750 | Intraspecies and domain-domain interaction based | [87] |
| EEF1A1 | TB10.6 k15.2790 | Conserved in EEF1A1(Human)-US3(Human herpesvirus 1 Strain 17) | [58] |
| EEF1A1 | TB1927.2.5980 | Conserved in EEF1A1(Human)-CLP[Bacillus anthracis] | [86] |
| RPN1 | VP24 | Experimentally validated | [58,86] |
| RPN1 | DinB | Conserved in RPN1(Human)-DP04(Veronius pestis) | [86] |
| RPN1 | TB11.01.0040 | Intraspecies and domain-domain interaction based | [87] |
| RPN1 | TB10.61.2630 | Intraspecies and domain-domain interaction based | [87] |
| RPN1 | TB1927.8.4890 | Intraspecies and domain-domain interaction based | [87] |
| SMARC2 | VP24 | Experimentally validated | [58] |
| SMARC2 | HemiH | Conserved in SMARC2(Human)-HemiH(Bacillus anthracis) | [86] |
| SMARC2 | GabD | Conserved in SMARC2(Human)-GabD(Bacillus anthracis) | [86] |
| SMARC2 | TB10.70.0630 | Intraspecies and domain-domain interaction based | [87] |
| TUBB | VP24 | Experimentally validated | [58] |
| TUBB | SpeA | Conserved in TUBB(Human)-SpeA(Veronius pestis) | [86] |
| TUBB | TB10.6 k15.2790 | Conserved in TUBB(Human)-US3(Human herpesvirus 1 Strain 17) | [58] |
associated proteins (Opa) and type IV pili, leads to poor immune control of NM infections [73]. Common to NM and Tb diseases is also the depletion of the complement cascade function. In Try-
panosoma infections by Tb gambiense and Tb rhodesiense, reduction in complement activity is caused by their massive activation of complement [74]. Human infection by NM begins by the colonization of the mucosal epithelium in cervix, conjunctiva, fallopian tubes, nasopharynx, rectum and urethra, by bacterial type IV pilin and Opa proteins [75] targeting host cell specialized adhesins.

4.1.3. Conserved host and pathogen-specific pathways

Our census shows that the three evolutionary distant pathogens develop similar strategies for host mitigation, defense and evasion by interaction with the same host proteins (Table 1). Detailed shared and unique PPIs occurring in different compartments are listed in Table 1. Host interactions with pathogen proteins happen in six different cellular compartments (green nodes). NM and Tb target both human immunoglobulin heavy constant gamma-1 (IGHG1) chain, tubulointerstitial nephritis antigen-like protein (TINAGL1), latent-transforming growth factor beta-binding protein-4 (LTBP4) and interleukin-18-binding protein (IL18BP). Host plasma membrane proteins include HLAs (human leukocyte antigen; HLA-B, HLA-DRA, HLA-DRB1 and HLA-DRB5), CD surface protein (CD74). Macrophage-expressed gene 1 protein (MPEG1), immunoglobulin superfamily member-6 (IGSF6) and sialic acid-binding Ig-like lectin-11 (SIGLEC11).

Cytoplasmic host targets for NM and Tb include apoptotic and inflammatory p53 and transforming growth factor beta-1-induced transcript 1 protein (TGFBI11), which regulates integrin, Wnt and TGFβ signaling pathways. Shared host targets are also HLA molecules HLA-DRB1 and HLA-DRB5 in Golgi apparatus and the endoplasmic reticulum. Shared cyto-nuclear interactions include proteasome related proteins (PSM2 and PSM3), transforming growth factor-beta regulator 1 (TBRG1) and Pre-B-cell leu-
kemia transcription factor 2 (PBX2).

4.2. Infection biology of the three pathogen networks: Specific and general targeting of host proteins

The interactions of the four conserved core proteins (Fig. 5) are contrasted by kingdom-specific interactions: eukaryotic try-
pansomes target host nuclear or the mitochondrial proteins, while prokaryotic bacteria target often also metabolic enzymes as well as host cell molecules by surface proteins [76], e.g. NM tar-
gets host microtubules [77]. Virus-interactions manipulate the host translation or transcription machinery as is the case for PCNA located in the cytoskeleton.

Many protozoan parasites adequately control growth and dif-
ferentiation processes for survival in the different environments they encounter during their life cycle. One way to do so may be by expressing different sets of mitogen-activated (MAP) kinases for each proliferative stage. For example, a Tb protein kinase con-
taining the signature of extracellular-signal-regulated kinases has been cloned and analyzed. Deletion of Tb MAPK2 results in delayed differentiation and growth inhibition and cell cycle arrest. Tb MAPK2 may thus control the growth of procyclic form try-
pansomes in the tsetse midgut [78]. A second MAP kinase, Tb MAPK5, seems to have a regulatory function during bloodstream form differentiation of Tb. As a result, knockout parasites seem to express an increased sensitivity to the stumpy induction factor, and knockout parasites differentiated prematurely in mice and cul-
ture [78]. Whether or not a physical interaction exists between any of these MAP kinases and host elongation factor 1-alpha 1 and/or tubulin-5 beta chain needs to be confirmed. The potential physical interaction might relate to the switch on/off of differentiation pro-
cesses that occur during the blood stage, and/or to guide parasite attachment to the host cell for example during parasite migration to the brain.

Additionally, the human oligosaccharyltransferase, involved in posttranslational protein modification, is localized in the endoplas-
mic reticulum [88], suggesting a similar localization with Tb puta-
tive proteins involved in protein folding and protein transport from the endoplasmic reticulum. Moreover, in Tb, a polymerase K (POLK) gene is present in 10 copies, in tandem repeats, and is not subjected to alternative splicing [89]. In T. cruzi, two copies of the POLK gene with differences in their encoded protein sequences outside of their catalytic core regions have been described. TcPOLK overexpression increases T. cruzi survival when exposed to H2O2 treatment. Phylogenetic analysis of TcPOLK copies together with its orthologues shows that the Polk of trypanoso-
matids has diverged early from other eukaryotes counterparts and at least one of the copies is located at the parasite mitochondrion and supports in vitro DNA synthesis [79] probably involved in error-prone translesion synthesis. Whether or not the predicted Tb POLK described above is involved in error prone translesion synthesis need to be explored as well as its physical interaction with the human dolichyl-diphosphooligosaccharide-protein glyco-
syltransferase subunit 1.

In Tb, exit from the endoplasmic reticulum is mediated by coat protein II (COPII)-coated vesicles that bud from distinct ribosome-
free regions of the ER membrane known as ER exit sites (ERES). Tb success relies as an infecting pathogen on coat protection by the variant surface glycoprotein (VSG). VSG is membrane-bound by a glycolipid (GPI) anchor, attached in the earliest compartment of the secretory pathway, the endoplasmic reticulum. Therefore, its exit from the endoplasmic reticulum is required for its proper expression at the surface of the parasite [80].

The Venn Diagram summarizes all these interactions (Fig. 3). A high proportion of overlap exists between Neisseria- and trypanosome-human interactions, and these conserved PPIs focus on host proteins related to immune system signaling, response to stress and reprogramming of host cell senescence. On the contrary, overlapping interactions between ZE and NM focus on mRNA sta-

4.3. Validation of the conserved four proteins targeted by pathogens

To further validate the four proteins commonly targeted by NM, TB, ZE, all available experimental evidence for these proteins was collected (Table 3), notably for ZE there is extensive direct experimental validation, for NM and Tb we collected circumstantial evi-
dence. Nevertheless, this still could be a rare coincidence from the particular pathogen choice. By three different large-scale data sets we hence did next confirm that the four human proteins found to be targeted comparing the evolutionary very distant pathogens NM, TB, ZE are no chance observation but these four human pro-
teins are very often targeted by human pathogens. Specifically, we found evidence from well-curated databases with experimental data that
(i) 56 human pathogenic bacteria and viruses target these four proteins,
(ii) the well researched new pandemic pathogen SARS-CoV-2 targets two of these four human proteins and
(iii) nine human pathogenic fungi (yet another evolutionary distant organism group) target three of the conserved proteins by 130 high confidence interactions.

Together, all these data confirm that these four central proteins are highly conserved and hence important proteins that are targeted by many human pathogens including bacteria, fungi, viridae and trypanosomes.

4.4. Four proteins are highly conserved targets as they are targeted by many human pathogens

The clade of only four interacting proteins present in the host and very strongly and conserved targeted seems to assure the proper decoding, produce cellular proteins and relocate them to the appropriate place. All four are important regulators of the cytoskeleton as explained in the following: EF1α is undoubtedly important for protein translation. However, EF1α proteins also modulate cytoskeleton activities, exhibit chaperone-like activity, and are key proteins for cell proliferation and cell death in human tumours [81]; EF1α assists replication of many RNA viruses, such as the respiratory syncytialvirus (RSV). In this case, down-regulation of EF1α restricts the expression of viral genomic RNA and the release of infectious virus [82]; in parasites, EF1α has also been implicated in pathogenesis [83] and host cell invasion [84]. The other three proteins also organize the cytoskeleton and we think that cytoskeletal signaling and its organization are both critical for the host defense as well as the success of the pathogen, and hence this results in these highly conserved targets: SWI/SNF complex subunit SMARCC2 is the matrix-associated actin-dependent regulator of chromatin subfamily C. It is targeted by bacteria and viridae but not targeted by fungi. The dolichyl-diphosphooligosaccharide–protein glycosyltransferase subunit 1 (RPN1) transfers glycosides to phospholipids and oligosaccharides. This function is again important in the cytoskeleton. Finally, the tubulin beta-5 chain (TUBB) is an important component of the cytoskeleton. The central functions of the four proteins (Supplementary File 8c) support the idea that many pathogens profit from targeting central host processes via the cytoskeleton and host entry by targeting these four proteins. Hence, this descriptive and in silico finding of high conservation of a human protein clade in infection requires experimental follow up to better understand the exact function of the four proteins in human-pathogen interactions and for the infection process. Ultimately, this could open up new therapeutic intervention strategies targeting host proteins pharmacologically which has in addition the promise of preventing fast resistance mutations which evolve from survival selection pressure by targeting pathogen proteins (e.g. antibiotic resistance of numerous pathogens Supplementary File 8c).

5. Conclusions

Septic pathogens Trypanosoma, Neisseria and Ebola virus invade the host in different manners; however, we found four common human proteins which are targeted by each of the pathogens. Experimentally validated direct interactions are available for VP24 from ZE to all four conserved human proteins and there is circumstantial evidence for NM and Tb. Next, we systematically looked at all available human-pathogen database data comparing human pathogenic bacteria, viruses and fungi. We found that indeed these four proteins are often targeted by human pathogens according to the available database information. This includes confirmation by experimental data in several specific cases including two of these proteins to be targeted by SARS-CoV-2. Our results suggest an unexpected human protein clade of four proteins targeted often by human pathogens: EF1α, involved in host cell invasion; tubulin beta-5 chain; cytoskeletal glycosyltransferase RPN1; actin-dependent regulator SMARCC2. All four proteins are involved in cytoskeletal organization and by this in further central human host processes.

Credit authorship contribution statement

Conceptualization: APS, SG, TD; Data curation and Formal analysis: All; Funding acquisition: APS, TD; Investigation: All; Methodology: SG, EB; Project administration: TD; Software: SG; Supervisions: APS, TD; Validation: All; Visualization: SG, EB; Writing - original draft: APS, SG, TD; Writing - review and editing: All.

Ethics approval and consent to participate or to publish

Not applicable, this manuscript does not include human participants, human data or human tissue.

Availability of data and material

The datasets generated and analysed during the current study are all available in the manuscript and its supplementary materials.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

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