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Virus–host interactomes — antiviral drug discovery
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One of the key questions in virology is how viruses, encoding relatively few genes, gain temporary or constant control over their hosts. To understand pathogenicity of a virus it is important to gain knowledge on the function of the individual viral proteins in the host cell, on their interactions with viral and cellular proteins and on the consequences of these interactions on cellular signaling pathways. A combination of transcriptomics, proteomics, high-throughput technologies and the bioinformatical analysis of the respective data help to elucidate specific cellular antiviral drug target candidates. In addition, viral and human interactome analyses indicate that different viruses target common, central human proteins for entering cellular signaling pathways and machineries which might constitute powerful broad-spectrum antiviral targets.

Experimental methods to study virus–host interactomes
Proteomic changes can be studied at the level of individual virus–host protein interactions, organelles, and whole cells. 2D-polyacrylamide gel electrophoresis and mass spectrometry (MS) have been used in numerous studies to identify gene expression patterns in infected cells [1**]. While 2D-PAGE methods and variations thereof allow the identification, relative quantification, and comparison of the abundant protein in infected versus noninfected samples, very small, very large, hydrophobic and low-abundance proteins are difficult to resolve. Drawbacks are a frequent limitation in reproducibility and lack of HT capabilities. Mass spectrometry is an excellent means of protein identification. Isobaric tagging techniques [2] including stable isotope labeling by amino acids (SILAC) in cell culture allow quantitative examination of virus–host cell relations and have improved the field significantly [3*]. MS-based techniques have recently also been used to characterize host–pathogen interactions within purified, mature virus particles including vaccinia virus [4], influenza virus [5], HIV [6], vesicular stomatitis virus [7], SARS-Corona virus [8*], and several herpes viruses [9*]. Finally, the two-hybrid-based mammalian PPI trap (MAPPIT), which drives a cellular signaling cascade with an endogenous transcriptional reporter, has been used as a HT assay for HCV and HIV-1 proteins and for human protein interactome analysis [10,11].

Yeast two-hybrid (Y2H) is a popular and intensely used alternative for studying virus–host protein–protein interaction on a global proteome scale [12*]. It exploits yeast genetic engineering and the modularity of eukaryotic transcription factors (TFs) utilizing DNA-binding
domains (DBD) and activating domains (AD). Bait proteins fused to DBD and prey proteins fused to AD reconstitute functional TFs (e.g., yeast GAL4) upon interaction of the fusion proteins to be tested. The system can be used in a HT manner performing array-based matrix screens (e.g., all virus proteins against each other) or screens of individual bait virus proteins against cDNA prey libraries of different origin [13\textsuperscript{*}]. Former allows the establishment of intraviral [14\textsuperscript{*}], latter of virus–host interactomes [15\textsuperscript{**},16]. Concepts of virus–host interactome studies are outlined in Figure 1. There are limitations to the Y2H system as the transcriptional reporter system is on the basis of nuclear localization, which limits the analysis of hydrophobic membrane proteins because of possible disruption of the nuclear membrane. Also, the lack of mammalian translational modifications might contribute to the detection rate of about one quarter of all interactions as estimated in recent studies [17\textsuperscript{**},18]. False positive interactions, a further drawback, can be efficiently prevented using proper controls, thus resulting in high-quality large-scale screens [19]. Increasing reliability by decreasing false-negative rates has recently been achieved by systematically combining screening strategies using novel N-terminal bait and C-terminal bait and prey fusion-protein vectors [20\textsuperscript{*}]. Biological significance can be addressed by the validation of Y2H interactions with different biochemical and/or mammalian cell-based methods. Co-immunoprecipitation was successfully applied to co-produced proteins containing short fusion tags in mammalian cells [14\textsuperscript{*},21]. As this method is very time-consuming, it is only applicable to small-scale evaluations. The LUMIER (Luminescence-based Mammalian IntEractome mapping) [22\textsuperscript{*}] assay or modified versions thereof [15\textsuperscript{**}] are pull-down assays with bait proteins used as fusions to Flag-tag or Protein A and with prey proteins used as fusions to luciferase (and vice versa) for detection. This methods as well as protein fragment complementation assays (PCAs) such as split-YFP-based or split-luciferase-based methods are well suited for HT screening. In the latter two assays, the test proteins are fused to subdomains of YFP [23\textsuperscript{*}] or luciferase [24]. Upon physical interaction of the bait and prey proteins, the respective YFP or luciferase fragments reconstitute the fluorescent or enzymatic activities, respectively. A split beta-lactamase interaction assay has very recently been described as a cell-free test for the screening of small molecular peptide inhibitors [25]. A very intriguing fluorescent two-hybrid (F2H) assay allows the direct visualization and analysis of PPIs in single living cells utilizing a stable nuclear interaction platform [26\textsuperscript{*}].
Identification of cellular antiviral drug target candidates by HT screening

In recent years, there have been several publications on genome-wide genetic screens for host-cellular cofactors of influenza virus, hepatitis C virus (HCV), human immunodeficiency virus (HIV), severe acute respiratory virus (SARS-CoV), and other viruses [12*,15**,27–35,36**].

In the field of influenza virus, dsRNA, and siRNA, yeast-2-hybrid screening was applied to select host genes crucial for influenza virus infection [27,31–33,35]. Shaw et al. compared the results of these screens and sorted the major selected cellular cofactors into six functional categories [36**]. The gene classification clusters are ribosome, COPI vesicle, proton-transporting V-type ATPase complex, spliceosome, nuclear pore/envelope, and kinase/signaling. The involvement of ribosomes is not surprising, as they are responsible for catalyzing synthesis of proteins from amino acids, not only of host cells but also of viruses. In addition to influenza virus, the ribosome has been demonstrated to play roles during infection by other viruses. For example, SARS-CoV nonstructural protein (Nsp-1) binds to the 40S ribosomal subunit to inactivate the translational activity of these subunits and induces host-mRNA degradation in SARS-CoV-infected cells [37,38]. The vacuolar-type H+-ATPase (vATPase) complex acidifies endosomes. This step is required for the fusion of the viral and endosomal membranes and results in subsequent release of the viral genome into the cytosol. Thereby, the activity of the vATPase complex is theoretically important for the entry of all viruses taking advantage of the host cell’s endocytic machinery, such as SARS-CoV [39], and semiliki forest virus [40], but perhaps not for the viruses which enter cells via direct membrane fusion. The importance of nuclear pore/envelope-associated proteins has been indicated previously [12*]. For instance, human KPNA1 (karyopherin alpha 1 or importin alpha 5) is an interacting partner of SARS-CoV accessory protein 6 and HCV nonstructural protein NS3 [34,41]. Human KPNA2 (karyopherin alpha 2 or importin alpha 1) interacts with multiple viral pathogen groups including SARS-CoV, adenovirus, and HIV [41,42*]. Human KPNB1 (karyopherin beta 1), which forms a heterodimer with karyopherin alpha to assist the import of proteins with a nuclear localization signal (NLS), was found in all of the five influenza screens [36**]. Kinase signaling is a very broad category. In this category, the mitogen-activated protein kinase (MAPK) signaling seems to play an important role during not only influenza virus infection but also HCV infection [34,36**]. In addition, the phosphorylation-mediated JAK/STAT signaling regulating expression of interferon-stimulated genes is also involved in response to multiple viral pathogens [42*]. The abundant kinase signaling category provides many druggable targets such as MAP2K3 and CDC-like kinase 1, inhibition of which reduced influenza virus replication by more than two orders of magnitude [32].

Bioinformatical identification of cellular targets as broad-range antiviral targets

Naturally, drugs targeting viral proteins tend to be virus-specific. Drugs directed against cellular proteins or signaling pathways potentially have a much broader spectrum of antiviral activity, as different viruses may require similar cellular functions for replication. As described before, several druggable cellular proteins might serve as antiviral targets. For various viruses, especially for influenza virus, systematic surveys of small-compound libraries for antiviral activities were performed utilizing highly sophisticated HTS methodologies. However, as in the case of influenza virus, no efficient antivirals have been approved this way until now [43].

Instead of determining gene expression by measuring mRNA levels in virus-infected cells, it might be more efficient to look for direct PPIs utilizing, for example ‘classical’ Y2H screening in combination with HT. Several intraviral protein networks (Epstein–Barr Virus [EBV], Influenza A Virus [FLUAV], HCV, Herpes-Simplex-Virus 1, Kaposi’s Sarcoma-associated Herpesvirus, SARS-CoV, and Varicella Zoster Virus) and virus-host protein networks (Dengue Virus, EBV, FLUAV, HCV, Vaccinia Virus) were bioinformatically analyzed indicating that viral proteins target highly central human proteins which define Achilles’ heels of the human interactome [44**]. Accordingly, viruses seem to share host proteins as targets for overtaking cellular signaling pathways and machineries which might constitute attractive broad-spectrum antiviral targets.

A SARS-CoV HTY2H screen identifies immunophilins as broad-spectrum anticonviral targets

A HTY2H approach was very recently used successfully in the coronavirus field: in unbiased, hypothesis-free screens, the SARS-CoVorfome including subfragments thereof were tested against human cDNA libraries leading to the identification of human immunophilin (cyclophilins and FK506-binding proteins [FKBP]) gene families as prerequisites for virus replication. The respective proteins have been known for many years to be inhibited by the immunosuppressive compounds cyclosporine A (CsA) and FK506. The interaction of the SARS-CoV Nsp1 protein with the cyclophilins PPIA, PPIG, and PPIH, with the FKBP1 FKBP1A (FKBP12) and FKBP1B (FKBP12.6) as well as with the calcipressin RCAN1 and RCAN3 (also confirmed by a modified luminescence-based mammalian interactome-mapping assay) [22**] was eye-catching, particularly with regard to their influences on the cellular serine phosphatase calcineurin (Cn) [15**]. On the basis of the well-known complex formations between cyclophilins/ CsA and FKBP/FK506, two important issues arose on the pathogenesis and on the replication of SARS-CoV. Figure 2 shows the influences of cyclophilin/CsA and FKBP/FK506 and the possible involvement of SARS-CoV Nsp1 on
NFAT regulation and virus replication. Regarding pathogenicity, the induction of the immunologically very important Cn/NFAT (Nuclear Factor of Activated T-cells) pathway by the virus and the viral Nsp1 might provide a possible explanation for the highly disordered cytokine levels (‘cytokine storms’) in SARS patients [45]. However, it is not known how the Nsp1 protein influences NFAT induction. It was further shown that CsA inhibits replication of not only SARS-CoV, but also of two other human coronaviruses, HCoV-NL63 and HCoV-229E, and of the animal CoVs Feline CoV (FCoV), Transmissible Gastroenteritis Virus (TGEV, pig), and Infectious Bronchitis Virus (IBV, bird). As CsA inhibits representatives of all three CoV genera (alpha-CoV, beta-CoV, gamma-CoV) it might thus serve as a pan-coronavirus inhibitor. As gammacoronaviruses lack Nsp1, it is highly possible that other coronaviral proteins are also involved in the interaction with cyclophilin. Such an interaction has been demonstrated in vitro for the SARS-CoV nucleocapsid protein [46].

The mechanism of inhibition of coronavirus replication by CsA is not clear. It is reasonable to assume, however, that in analogy to HIV-1 [47] and HCV [48], the peptidyl-cis-trans isomerase and chaperon activities of the cyclophilins are blocked by CsA, leading to improper folding of viral proteins. In case of HIV-1, a crystal structure of the complex between cyclophilin and the viral capsid protein
CsA also exerts inhibitory effects on HSV-1 [52], vaccinia virus [53], BK polyoma virus [54] and influenza virus by cyclophilin A-dependent and cyclophilin A-independent pathways [55]. FK506 also inhibits the human CoVs SARS-CoV, NL63, and 229E [56*]. It was found to have an inhibitory effect on chronically but not on newly HIV-1-infected cells (up to 10 μg/ml FK506 [57]). Replication of replicon HCV RNA up to concentrations of 3 μg/ml was also not affected [48]. Orthopoxviruses however were inhibited by FK506 [58]. Anti-immunophilin drugs are very promising antiviral candidates. However, severe immunosuppressive side effects are normally undesirable. These can be overcome by the development of nonimmunosuppressive derivatives of the drugs which still display antiviral activity by destroying the calcineurin-binding site and the preservation of the cyclophilin-binding activity. A very promising example is the CsA derivative Debio-025, SCY-635, NIM-811 which has been demonstrated to inhibit HCV and HIV-1 replication very efficiently [59**].

Conclusions

Integrated viral ORFeome databases designed to generate versatile collections of viral ORFs [60], viral [61], and virus–host protein [62] interaction databases are extremely valuable tools for the comparison and analysis of an increasing several virus interactomes as well as individual proteins at the intraviral and at the virus–host level. Several assays are available to study virus–host relations, each carrying intrinsic advantages and disadvantages. Techniques addressing interactions at the protein, not only at the RNA level, might be more promising with respect to inhibitor identification because of, for example the role played by post-transcriptional modifications. Indeed, by HTY2H screening of the SARS-CoV ORFeome against human cDNA libraries, we have identified members of the cyclophilin and FKBP protein families as prerequisites for coronavirus replication. Interaction of the SARS-CoV Nsp1 protein with these cellular proteins led to the discovery of the natural cyclophilin A ligand CsA as an inhibitor of animal and human coronaviruses (pan-coronavirus inhibitor) and of the natural FKBP-ligand FK506 as an inhibitor of human coronaviruses. A second but independent aspect of this interaction was the discovery of the upregulation of the immunologically very important Ca/NFAT signaling pathway by SARS-CoV Nsp1 and the virus itself, which might provide an explanation of the so-called cytokine storms in SARS patients. From a technical point of view, it may be noted that the described Nsp1 interactions occurred with low abundance in the Y2H screens. The relevance of these cellular protein families to coronavirus biology was first realized by very careful ‘eye inspection’ of and contemplation on the interaction data keeping related literature in mind. That is, the construction of large and colorful interaction maps per se does not necessarily give hints on relevant cellular target molecules or signaling pathways for antiviral therapy. Undoubtedly, interaction databases and databases specializing in drug target PPIs, PPI-inhibiting small molecules, and integrative systems for assessing the druggability of PPIs [63–65] will provide invaluable computerized tools for the development of antivirals in the future.

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