Hemorrhagic fevers caused by arenaviruses are among the most devastating emerging human diseases. Considering the number of individuals affected, the current lack of a licensed vaccine, and the limited therapeutic options, arenaviruses are arguably among the most neglected tropical pathogens and the development of efficacious anti-arenaviral drugs is of high priority. Over the past years significant efforts have been undertaken to identify novel potent inhibitors of arenavirus infection. High throughput screening of small molecule libraries employing pseudotype platforms led to the discovery of several potent and broadly active inhibitors of arenavirus cell entry that are effective against the major hemorrhagic arenaviruses. Mechanistic studies revealed that these novel entry inhibitors block arenavirus membrane fusion and provided novel insights into the unusual mechanism of this process. The success of these approaches highlights the power of small molecule screens in antiviral drug discovery and establishes arenavirus membrane fusion as a robust drug target. These broad screenings have been complemented by strategies targeting cellular factors involved in productive arenavirus infection. Approaches targeting the cellular protease implicated in maturation of the fusion-active viral envelope glycoprotein identified the proteolytic processing of the arenavirus glycoprotein precursor as a novel and promising target for anti-arenaviral strategies.

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Introduction

Arenaviruses are important emerging human pathogens

Several arenaviruses cause severe viral hemorrhagic fevers (VHF) in humans and represent a serious public health problem (Geisbert and Jahrling, 2004). Lassa virus (LASV) in Africa causes several hundred thousand infections per year resulting in significant mortality and morbidity (McCormick and Fisher-Hoch, 2002). On the South American continent, the arenaviruses Junin (JUNV), Machupo (MACV), Guanarito (GTOV), and Sabia virus (SABV) have emerged as etiological agents of severe VHF in Argentina, Venezuela, Bolivia, and Brazil, respectively.
The worldwide-distributed prototypic arenavirus lymphocytic choriomeningitis virus (LCMV) is also a neglected human pathogen of clinical significance, especially in pediatric medicine (Barton et al., 2002) and represents a threat to immuno-compromised individuals, as tragically illustrated by recent fatal cases of transplant-associated LCMV (Barton et al., 2002) and represents a threat to immuno-compromised virus lymphocytic choriomeningitis virus (LCMV) is also a neglected (Buchmeier et al., 2007). The worldwide-distributed prototypic arena-164

predicted hairpin structure (Fig. 1). The S RNA encodes the viral orientation, separated by a non-coding intergenic region (IGR) with a strategy to direct the synthesis of two polypeptides in opposite enveloped viruses with a bisegmented negative strand RNA genome and only a short summary will be given here. Arenaviruses are excellent recent reviews (Buchmeier et al., 2007; de la Torre, 2009) humans and will be covered in the present review.

of novel drugs targeting different steps in the arenavirus life cycle is responses, to control, and ultimately clear the virus. The development of viral replication and spread may provide the patient's immune system a window of opportunity to develop anti-viral immune responses, to control, and ultimately clear the virus. The development of novel drugs targeting different steps in the arenavirus life cycle is therefore a promising strategy to combat arenavirus infection in humans and will be covered in the present review.

Molecular and cell biology of arenaviruses

The molecular and cell biology of arenaviruses has been covered by excellent recent reviews (Buchmeier et al., 2007; de la Torre, 2009) and only a short summary will be given here. Arenaviruses are enveloped viruses with a bisegmented negative strand RNA genome and a life cycle restricted to the cytoplasm. Each genomic RNA segment L (ca 7.3 kb) and S (ca 3.5 kb) uses an ambisense coding strategy to direct the synthesis of two polypeptides in opposite orientation, separated by a non-coding intergenic region (IGR) with a predicted hairpin structure (Fig. 1). The S RNA encodes the viral glycoprotein precursor, GPC, and the nucleoprotein, NP, (ca 63 kDa), whereas the L RNA encodes the viral RNA dependent RNA polymerase (RdRp, or L polymerase) (ca 200 kDa), and a small RING finger protein Z (ca 11 kDa). Arenavirus GPC is synthesized as a single polypeptide chain (ca 75 kDa) and post-translationally cleaved by the cellular procoprotein convertase (PC) subtilisin kexin isoizyme-1 (SKI-1/site-1 protease (SIP) to yield the mature virion glycoproteins GP1 (40–46 kDa) and GP2 (35 kDa) (Beyer et al., 2003; Lenz et al., 2001; Pinschewer et al., 2003; Rojek et al., 2008a). The GP1 part mediates virus interaction with host cell surface receptors and is located at the top of the mature GP spike present in the viral envelope. GP1 is associated via ionic interactions with the transmembrane GP2 that forms the stalk of the spike. Arenavirus GP2 resembles the fusion-active membrane-proximal parts of other enveloped viruses. The cellular receptor for LASV and most isolates of LCMV is α-dystroglycan (α-DG), a cell surface receptor for proteins of the extracellular matrix (ECM) (Cao et al., 1998). The New World arenaviruses JUNV, MACV, GTOV, and SABV can use human transferrin receptor 1 (TfR1) as a cellular receptor (Radoszitzky et al., 2007). Upon attachment to the target cell, arenavirus particles are taken up by endocytosis and delivered to acidified endosomes where low pH induces membrane fusion (Borrow and Oldstone, 1994). New World arenaviruses like JUNV that use human TfR1 enter the cell via clathrin-mediated endocytosis (Martinez et al., 2007). In contrast, the Old World arenaviruses LASV and LCMV, which depend on α-DG, use a distinct and unusual pathway for cell entry that is independent of clathrin, caveolin, dynamin, and actin and bypasses classical routes of incoming early endosomal trafficking (Quirin et al., 2008; Rojek et al., 2008b,c).

Formation and release of arenavirus infectious progeny from infected cells by budding requires that assembled viral RNPs associate at the cell surface with membranes that are enriched in viral GPs. For many enveloped RNA viruses viral budding is mediated by a matrix (M) protein that acts as a bridge between the RNP and GP. In arenaviruses, the Z protein functions as the main driving force of arenavirus budding (Perez et al., 2003; Strecker et al., 2003) and the process is mediated by the Z proline-rich late (L) domain motifs (PTAP and PPPP) known to control budding of several other viruses via interaction with specific host cell proteins (Freed, 2002).

Current drugs to combat arenaviruses

The only licensed drug for treatment of human arenavirus infection is the nucleoside analogue ribavirin (Rib) (1-β-D-ribofuranosyl-1,2,4-triazole-3-carboxamide) (Parker, 2005). In vitro and in vivo studies have documented the prophylactic and therapeutic value of Rib against several arenaviruses. Rib reduced both morbidity and mortality in humans associated with LASV infection (McCormick et al., 1986), and experimentally in MACV (Kilgore et al., 1995) and JUNV (Weissenbacher et al., 1987) infections, if given early in the course of clinical disease. The mechanism of action of Rib against arenaviruses is currently not entirely clear. Studies in the LCMV model revealed that the anti-viral effect of Rib is not associated with significant increases in virus mutation frequencies, but has a dramatic effect on viral RNA synthesis (Ruiz-Jarabo et al., 2003) Over the past two decades efforts have been made to discover novel drug candidates to combat arenaviruses and only a short and by no means comprehensive overview can be given here: inhibitors of IMP dehydrogenase (Andrei and De Clercq, 1993), the S-adenosylhomocysteine (SAH) hydrolase (Andrei and De Clercq, 1990), phenothiazine compounds (Candurra et al., 1996), brassinosteroids (Wachsmann et al., 2000) and myristic acid (Cordo et al., 1999) have been reported to have anti-arenaviral activity. Several zinc-finger-reactive compounds with antiretroviral potential showed activity against arenaviruses (Garcia et al., 2000, 2006) and evidence has been provided for an involvement of the viral Z protein in their mechanism of action (Garcia et al., 2010). The pyrazine derivative T-705 (6-fluoro-3-hydroxy-2-pyrazinecarboxamide) showed activity against the arenaviruses JUNV, Pichinde (PICV), and Tacaribe virus (TACV) in vitro and protected against PICV infection in a hamster model (Gowen et al., 2007), with efficacy in late stage infection (Gowen et al., 2008).

Early attempts to target arenavirus entry employed sulfated polysaccharides (Andrei and De Clercq, 1990) and revealed inhibitory effects of charged polymers on arenavirus infection, however, the exact mechanism of action remained elusive. A more recent study evaluated phosphorothioate oligonucleotides in the LCMV model and revealed that amphipathic DNA polymers are potent inhibitors of arenavirus infection (Lee et al., 2008). The DNA polymers act at the level of cell entry and target the interaction between LCMVGP and its cellular receptor, α-DG, without affecting later steps in replication. While these studies are of great value as proof-of-principle, to the best of our knowledge none of these drug candidates has undergone further evaluation in pre-clinical and clinical trials.

Discovery of novel inhibitors of arenavirus entry by small molecule screening

Since entry into the host cell is the first step of every virus infection it represents a promising target for attacking the virus before it can gain control over the host cell machinery for replication. In recent years, viral entry inhibitors have emerged as a new class of anti-viral drugs, with enfuvirtide being successfully used for clinical treatment of humans

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infected with human immunodeficiency virus (HIV) (Este and Telenti, 2007; Qian et al., 2009) and potent small molecule inhibitors for entry of other viruses, including influenza (Vanderlinden et al., 2010) and hepatitis C virus (Baldick et al., 2010) have been reported.

Considering the limited public health infrastructure in endemic regions, highest priority is given to the development of novel synthetic drugs against human pathogenic arenaviruses that can be produced at low cost, delivered orally, and can be stored in tropical climates. To achieve this goal, several attempts at drug discovery based on screening large collections of synthetic small molecules have been undertaken. High throughput screening (HTS) of small molecular libraries has emerged as a major source of potent agonists and antagonists for a great number of molecular interaction implicated in various biochemical processes (Boger et al., 2003). Despite the possible involvement of several molecular interfaces, a large number of studies on protein–protein interaction indicate that in most cases, small clusters of amino acid residues mediate the energetically most important interactions (Livnah et al., 1996; Wells, 1996; Wells and de Vos, 1996; Wrighton et al., 1996) that may be perturbed by small synthetic molecules. Libraries of small molecules can therefore be used to screen for and identify molecules that promote protein–protein interactions (agonists) or inhibit protein–protein interactions (antagonists) (Boger et al., 1998).

A first attempt using HTS of small molecules to identify inhibitors of arenavirus infection employed a virus-induced cytopathic effect (CPE)-based assay involving the non-pathogenic New World arenavirus TACV. HTS of a random library identified a potent small molecule inhibitor of TACV and several other New World arenaviruses, ST-294 (Fig. 2) (Bolken et al., 2006), providing proof-of-principle for the feasibility of this type of approach.

All hemorrhagic arenaviruses are classified as biosafety level (BSL)-4 pathogens requiring high security containment laboratories for work with live viruses. In order to identify specific inhibitors of cell entry of hemorrhagic arenaviruses, subsequent drug screening strategies made use of recombinant retroviruses bearing the GP of hemorrhagic arenaviruses in their envelope and a luciferase reporter in their genome (Larson et al., 2008; Lee et al., 2008). Since arenavirus cell attachment and entry are mediated exclusively by the envelope GP, these pseudotype platforms allow rapid and reliable screening of small molecule libraries in a HTS format. On one study, pre-selected combinatorial small molecule libraries were chosen instead of random libraries, allowing a significant reduction of the scope of the screen to less than 100,000 compounds. Combinatorial chemical libraries generated by solution-phase synthetic techniques (Boger et al., 1998) have proven to be powerful sources of novel inhibitors or agonists of a variety of biochemical interactions. The use of this type of library has identified erythropoietin (EPO) mimetics (promote EPO receptor dimerization) (Berg et al., 2002), inhibitors of Myc/Max dimerization (Berg et al., 2002), inhibitors of LIF/IL-6 receptor dimerization (Boger et al., 2000), inhibitors of the binding of the protease MMP-2 to αvβ3 integrin (Boger et al., 2001; Silletti et al., 2001), and inhibitors of HIV protease (Chang et al., 2010). Positive screening of circa 80,000 compounds with LASV pseudotypes yielded initially circa 1% hits (Lee et al., 2008). Based on the combinatorial nature of the libraries, the hits tended to appear in clusters of structurally related compounds. Counter-screening employing pseudotypes of the unrelated vesicular stomatitis virus (VSV) yielded 32 single compounds and 53 mixtures of 4-to-10 compounds each that specifically blocked LASV GP-mediated infection. A series of leads were selected, including the candidate compounds 8C1 and 17C8 (Fig. 2). Candidate compounds 8C1 and 17C8 efficiently blocked infection of cells with live LASV, whereas candidate 17C8, but not 8C1, was also potent against JUNV and other New World hemorrhagic arenaviruses with comparable IC50 values. Mechanistic studies revealed that both 8C1 and 17C8 blocked the very last step of viral entry, pH-dependent membrane fusion. When tested in a cell-based arenavirus GP fusion assay, these compounds efficiently blocked membrane fusion mediated by the GPs of LASV and JUNV with an IC50 between 200 and 350 nM. The combinatorial nature of the chemical libraries allowed the identification of clusters of structurally related compounds with differential anti-viral activity, providing the basis for a first assessment of structure–activity–relationship (SAR). Subsequent examination of selected compounds in the same series as 17C8 and the related 16G8 (Fig. 2) identified several additional candidates that exhibited potent activity against LASV GP-mediated infection (Whitby et al., 2009). These compounds showed that the benzo[1,2-b:4,5-b′]dithiophene-2,5-dione moiety (X being either methyl- or methoxy-) impart the activity of the compound against the LASV GP, as
substitution at the third position around the central ring of various different-sized substituents, including methyl, benzyl, and phenethyl moieties, had no effect on the potency of the compound. Interestingly, after purification of the (R)- and (S)-isomers of 16G8, activity against the LASV pseudotypes was fully retained with (S)-16G8, whereas (R)-16G8 showed 15-fold lower activity. This indicates that the stereochemistry of the molecule is important for its function against arenavirus GP-mediated fusion. A similar effect was also observed with 17C8, with (S)-17C8 being significantly more potent than (R)-17C8 against LASV pseudotypes (Lee et al., unpublished results). The requirement for one stereoisomer over another suggests that our compounds bind to a specific structure on the GP to exert their effects, and efforts to identify where the compounds bind the GP and block fusion are currently underway.

Another screen for LASV entry inhibitors used a chemically diverse random library of 400,000 small molecule compounds and a lentivirus pseudotype platform (Larson et al., 2008). Applying a more stringent cut-off (>75% inhibition at 5 μM), HTS resulted in a hit rate of 1.2%. However, in contrast to the combinatorial library screen, where counter-screens reduced the number of hits by less than 50% (Lee et al., 2008), 90–95% of the hits in the random library screen were found to be not specific for LASV pseudotypes and/or cytotoxic (Lee et al., 2008). The most promising hit, a unique benzimidazole derivative, ST-37 (Fig. 2), exhibited an IC50 of 16 nM against LASV pseudotypes. Subsequent SAR and lead optimization yielded the compound ST-193 (Fig. 2) that inhibited pseudotypes of LASV, JUNV, MACV, GTOV, and SABV with IC50 of 2–12 nM (Larson et al., 2008). The candidate ST-193 was also potent against live viruses, supporting the use of pseudotype platforms for this type of screens. Mechanistic studies revealed that ST-193 and the formerly identified New World arenavirus entry inhibitor ST-294 inhibit GP-mediated membrane fusion (Larson et al., 2008; York et al., 2008). Interestingly, despite potent inhibition of cell entry of hemorrhagic arenaviruses by 17C8 and ST-193, both compounds seem to be far less active against LCMV, whose GP is structurally closely related to LASV (Larson et al., 2008; Lee et al., unpublished results). The resistance of LCMV GP towards ST-193 mapped to the C-terminal portion of GP2 and involves two residues in the transmembrane domain that show divergence between LCMV on the one hand and the hemorrhagic arenaviruses, including LASV, on the other hand (Larson et al., 2008). In sum, two independent small molecule screens using pseudotype platforms in combination with different library formats identified a set of broadly active potent inhibitors of arenavirus fusion. These studies provide proof-of-principle for this type of approach and establish arenavirus membrane fusion as a robust drug target to combat a broad range of arenaviruses. A notable strength of HTS of small molecule libraries is

![Fig. 2. Arenavirus entry inhibitors identified in small molecule HTS. (A) Arenavirus entry inhibitors identified by HTS of a random chemical library (Bolken et al., 2006; Larson et al., 2008). (B) Inhibitors of arenavirus entry obtained from HTS of combinatorial libraries (Lee et al., 2008). (C) SAR of the candidate compounds 17C8 and 16G8 obtained from the combinatorial library HTS (Lee et al., 2008). For details please see text.](image-url)
that no information on the structure of the viral envelope protein, receptor use, and pathway of endocytosis is needed. This approach holds therefore a great potential for the rapid discovery of cell entry inhibitors for newly emerging enveloped viruses that allow pseudotyping.

In addition to serving as promising leads for drug development against human pathogenic arenaviruses, the prototypic fusion inhibitors identified by these recent studies can also be used as molecular probes to dissect the mechanism of arenavirus membrane fusion. This is of particular interest in the context of arenaviruses as studies over the past years revealed unusual and unique features of arenavirus fusion that are not shared by other enveloped viruses. The GPs of arenaviruses show overall similarity to fusion-active class I envelope glycoproteins of other enveloped viruses (Eschli et al., 2006), but is unusual in that it contains a stable signal peptide (SSP) of remarkable length that is retained as an essential component of the fusion-active mature GP complex with the composition SSP/GP1/GP2 (Eichler et al., 2003; York et al., 2004). The arenavirus SSP contains 58 amino acids with two putative transmembrane domains and both the N- and the C-terminus in the cytosol and plays an important role in transport, maturation, and fusion-activity of the GP (Agnihotram et al., 2006, 2007; Saunders et al., 2007; York and Nunberg, 2006; York and Nunberg, 2007). In particular amino acid substitutions of the conserved residue K33 in the short ectodomain loop of SSP modulate the fusion pH of arenavirus GP, indicating an interaction between SSP and the fusion machinery of GP2. The New World arenavirus entry inhibitor ST-294 has been used in mechanistic studies to illuminate molecular details of JUNV fusion (York et al., 2008). Evidence was provided that ST-294 acts as a fusion inhibitor, targeting the interaction of the fusion-active GP2 subunit with SSP, and that amino acid substitutions at the residue K33 of SSP confer resistance to ST-294. Both ST-294 and the broadly active arenavirus fusion inhibitor ST-193 inhibit dissociation of GP1 from the SSP/GP1/GP2 complex induced by low pH, which occur concomitant with conformational changes in the fusion-active GP2 part (York et al., 2008).

The proteolytic processing of arenavirus glycoprotein precursor as a novel target for anti-viral therapy

A crucial step in the life cycle of arenaviruses is the proteolytic processing of the arenavirus GP precursor (GPC) by the cellular protease SKI-1/S1P, yielding fusion-active, mature GP. Processing of arenavirus GPC by SKI-1/S1P is essential for productive infection and viral spread (Beyer et al., 2003; Kunz et al., 2003; Lenz et al., 2001; Rojek et al., 2008a). In the host cell, SKI-1/S1P is involved in proteolytic processing of a defined set of cellular proteins, including the sterol regulatory element-binding proteins (SREBP-1 and SREBP-2), involved in lipid metabolism (Brown and Goldstein, 1997; Sakai et al., 1998), and the activating transcription factor 6 (ATF6), involved in the regulation of the cellular unfolded protein response (Schröder and Kaufman, 2005; Ye et al., 2000).

Interestingly, apart from the arenaviruses, SKI-1/S1P is also implicated in the proteolytic processing of the envelope GPs of the highly pathogenic Bunyavirus Crimean Congo HF virus (Vincent et al., 2003).

Efforts to develop specific inhibitors of SKI-1/S1P resulted in the design of recombinant serpins able to block SKI-1/S1P activity (Pullikotil et al., 2004). Expression of SKI-1/S1P-adapted ε1-antitrypsin variants efficiently inhibited the processing of LASV GPC and had a pronounced impact on production of infectious LASV from infected cells (Maïsa et al., 2009). Another study evaluated a cell-permeable, peptide-based inhibitor that combines a reactive chloromethylketone (CMK) moiety with peptides derived from the LASV GPC recognition motif of SKI-1/S1P, decanoyl (dec)-RRLL-CMK (Pasquato et al., 2006; Rojek et al., 2010). Dec-RRLL-CMK functions as a fast-acting suicide inhibitor of SKI-1/S1P (Pasquato et al., 2006) and showed significant anti-viral effects against the prototypic LCMV (Rojek et al., 2010). Off-target effects unrelated to SKI-1/S1P-mediated GPC processing were excluded employing a recombinant LCMV variant bearing a canonical furin recognition site (RRRR) in its GPC. Combination of the protease inhibitor with Rib resulted in additive drug effects. Remarkably, in cells deficient in SKI-1/S1P, the furin-dependent LCMV variant established persistent infection, whereas wild-type LCMV underwent extinction without emergence of SKI-1/S1P-independent escape variants. Together, these studies strongly suggested that inhibitors of SKI-1/S1P protease represent promising and powerful anti-arenaviral drug candidates.

The SKI-1/S1P-adapted serpins and the peptide inhibitor dec-RRLL-CMK used in these proof-of-principle studies both have limitations in their potential use as an anti-viral in vivo. Recently small molecule HTS performed by Pfizer Inc. identified a novel SKI-1/S1P inhibitor, the amino-pyrolidine amide compound PF-429242 (Hawkins et al., 2008; Hay et al., 2007). PF-429242 blocked SKI-1/S1P-mediated processing of the transcription factor SREBP in cell culture and treatment of mice with PF-429242 reduced expression levels of hepatic SREBP target genes and lowered rates of cholesterol and fatty acid synthesis (Hawkins et al., 2008). PF-429242 showed high stability, low toxicity and pharmacokinetic properties that made it an interesting drug candidate in the context of arenavirus infection. Indeed, PF-429242 efficiently prevented processing of GPC of LCMV and LASV, which correlated with the compound’s potent anti-viral activity against LCMV and LASV in cultured cells (Urata et al., 2011). In contrast, a recombinant LCMV expressing a GPC whose processing is mediated by furin was highly resistant to PF-429242 treatment. Mechanistically, PF-429242 did not affect virus RNA replication or budding, and had an only modest effect on virus cell entry, indicating that the anti-arenaviral activity of PF-429242 was mostly related to its ability to inhibit SKI-1/S1P-mediated GPC processing. These findings support the feasibility of using the small molecule SKI-1/S1P inhibitor PF-429242 as a novel anti-arenaviral drug.

Since SKI-1/S1P plays an important role in a number of physiological processes, including lipid metabolism and ER stress, the development of inhibitors that specifically target processing of arenavirus GPC while having minimal effects on the processing SKI-1/S1P’s cellular substrates has of course highest priority. Nevertheless, SKI-1/S1P inhibitors like PF-429242 are promising since hemorrhagic fevers caused by arenaviruses in humans are acute diseases with a relatively rapid course of infection (Geisbert and Jahrleing, 2004; McCormick and Fisher-Hoch, 2002). Consequently, drug intervention to treat hemorrhagic arenaviruses would be restricted to short periods of time. Dietary supplementation of cholesterol and other lipids to patients in intensive care appears feasible and would limit unwanted side-effects of PF-429242 on lipid metabolism. Moreover, cells deficient in SKI-1/S1P are still able mount a partial ER stress response (Ye et al., 2000), suggesting that short term inhibition of the SKI-1/S1P activity may not result in unacceptable levels of toxicity. Therefore, candidate drugs that lack absolute specificity and to some degree affect the processing of cellular targets of SKI-1/S1P may still represent important candidates to be evaluated as antiviral drugs.

Conclusions and outlook

Considering the significance of arenaviruses as important public health problems, the development of novel and potent anti-arenaviral drugs that can be manufactured at low costs remains an urgent need. Small molecule HTS performed over the past years identified novel classes of anti-viral drug candidates and provided proof-of-concept for the feasibility of this strategy for anti-arenaviral drug discovery. The successful identification of potent and specific inhibitors for arenavirus cell entry illustrates that small molecule HTS approaches for anti-viral drug discovery require neither structural information on the virus, nor knowledge on cellular factors involved. This type of approach is therefore highly suitable for rapid drug development against newly emerging human pathogenic viruses, provided their envelope proteins are amenable for pseudotyping. In addition to viral
entry, virtually every step of the arenavirus replication cycle could be targeted by this approach. The development of replicon systems for LASV (Hass et al., 2004), as well as a novel and powerful cell-based luciferase assay suitable for HTS of inhibitors of arenavirus budding (Capul and de la Torre, 2008) open the possibility to screen for compounds that act at additional steps in the viral life cycle. The availability of inhibitors against distinct steps of viral multiplication would open the possibility of combinatorial drug use, reducing the risk of emergence of drug-resistance.

The successful use of newly discovered arenavirus entry inhibitors highlights the potential of small molecule screens to provide novel small molecule agonists and antagonists that can be used as molecular probes to dissect biomolecular interactions. In particular the use of reversible small molecule inhibitors with mid- to low nanomolar probes to dissect biomolecular interactions. In particular the use of reversible small molecule inhibitors with mid- to low nanomolar probes to dissect biomolecular interactions. In particular the use of reversible small molecule inhibitors with mid- to low nanomolar probes to dissect biomolecular interactions.