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# Serpin functions in host-pathogen interactions

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Serpins are a broadly distributed superfamily of protease inhibitors that are present in all kingdoms of life. The acronym, serpin, is derived from their function as potent serine protease inhibitors. Early studies of serpins focused on their functions in haemostasis since modulating serine protease activities are essential for coagulation. Additional research has revealed that serpins function in infection and inflammation, by modulating serine and cysteine protease activities. The aim of this review is to summarize the accumulating findings and current understanding of the functions of serpins in host-pathogen interactions, serving as host defense proteins as well as pathogenic factors. We also discuss the potential crosstalk between host and pathogen serpins. We anticipate that future research will elucidate the therapeutic value of this novel target.

## 1 **Serpin Functions in Host-Pathogen Interactions**

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## 29 ABSTRACT

30 Serpins are a broadly distributed superfamily of protease inhibitors that are present in all  
31 kingdoms of life. The acronym, serpin, is derived from their function as potent serine proteases  
32 inhibitors. Early studies of serpins focused on their functions in haemostasis since modulating  
33 serine proteases activities are essential for coagulation. Additional research has revealed that  
34 serpins function in infection and inflammation, by modulating serine and cysteine proteases  
35 activities. The aim of this review is to summarize the accumulating findings and current  
36 understanding of the functions of serpins in host-pathogen interactions, serving as host defense  
37 proteins as well as pathogenic factors. We also discuss the potential crosstalk between host and  
38 pathogen serpins. We anticipate that future research will elucidate the therapeutic value of this  
39 novel target.

## 40 INTRODUCTION

41 Serpins are a superfamily of proteins. The family name is derived from the functional  
42 attributes of the members as they are serine protease inhibitors (serpin). They are the most  
43 broadly distributed protease inhibitors, and are present in all kingdoms of life including plants,  
44 animals, bacteria, archaea and viruses (*Silverman et al. 2001*). While the majority of serpins  
45 function as serine protease inhibitors, some serpins function as “cross-class” inhibitors of other  
46 kinds of proteases. For example, viral serpin CrmA inhibits a cysteine protease, interleukin-1  
47 beta converting enzyme (*Irving et al. 2002; Ray et al. 1992*). In addition, there are a few serpins  
48 that exhibit no inhibitory functions but participate in biological processes in other ways. For  
49 examples, serpin HSP47 serves as a chaperone, and ovalbumin (another serpin), functions as a  
50 storage protein (*Law et al. 2006*).

51 As potent protease inhibitors, serpins modulate a wide variety of proteolytic cascades thus  
52 controlling many physiological and pathological reactions. For instance, human serpins are  
53 found to regulate the proteolytic cascade that is central to blood clotting. Antithrombin, a serpin  
54 superfamily member, can inhibit multiple key enzymes in blood coagulation such as thrombin,  
55 activated factor X (FXa), FIXa and FXIa (*Aguila et al. 2017; Hepner & Karlaftis 2013*). In  
56 addition to blood coagulation, serpins also participate in a wide variety of other biological  
57 processes. These processes include thrombosis (*Van de Water et al. 2004*), immune-regulation  
58 (*Pemberton et al. 1988*), tumour-suppression (*Dzinic et al. 2017*), chromatin condensation  
59 (*Grigoryev & Woodcock 1998*) and apoptosis regulation (*Ray et al. 1992*). Furthermore, studies  
60 reveal serpins have clinical relevance. For example, patients with papillary thyroid cancer have  
61 high-concentrations of SERPINE2 and SLPI (secretory leukocyte protease inhibitor) (*Stein &*  
62 *Chothia 1991*). The serum concentrations of both anticoagulant proteins are considered markers  
63 for the development of this disease. Kallistatin, another serpin family member, has been shown  
64 to regulate cardiovascular function and blood vessel development. Its levels are elevated in  
65 patients with type 1 and type 2 diabetes with chronic diabetic complications (*Gateva et al. 2017*).

66 Recently, the study of serpin functions in infection and inflammation has been of particular  
67 interest, especially as more serpins from pathogens are identified and characterized. One  
68

69 example is crmA, a cowpox viral serpin, one of the smallest members of the serpin superfamily.  
70 CrmA is thought to be important in allowing viruses to avoid host inflammatory and apoptotic  
71 responses (Ekert *et al.* 1999; Renatus *et al.* 2000). Usually, viral genomes are compact to fit their  
72 unique life style. The presence of viral serpins indicates their essential functions for the survival  
73 of the pathogen and the infection of hosts. Since serpins are present in both pathogen and host  
74 organisms, we will discuss the functions of serpins on each side of these processes, and their  
75 potential interactions, in a variety of organisms during infection and inflammation.

76

## 77 SURVEY METHODOLOGY

78 In this study, we reviewed articles related to the functions of serpins in different organisms that  
79 either serve as hosts or pathogens. All references in this review were retrieved using search  
80 engines such as PubMed and Google Scholar. Keywords such as serpin, serine protease, host-  
81 pathogen interaction, infection and inflammation were used to search for the references. Figures  
82 related to protein structures were searched and modified from the Protein Data Bank.

83

## 84 Mechanism of Serpin Inhibition

85 The mechanism of serpin-inhibiting proteases relies on a reactive center loop (RCL) that  
86 interacts with the target enzyme. Structural studies reveal that serpins are characterized by a  
87 common core domain consisting of three  $\beta$ -sheets and 8-9  $\alpha$ -helices and a reactive center loop,  
88 as shown in Figure 1A (Moreira *et al.* 2014; Patschull *et al.* 2011; Pearce *et al.* 2008). Although  
89 this core region is present in all serpins, sequence homology among members in the family could  
90 be as low as 25%. A phylogenetic study of the superfamily divided serpins into 16 ‘clades’,  
91 designated clade A-P (Gettins 2002), and includes serpins from vertebrates, invertebrates, plants  
92 and viruses. A phylogenetic tree of representative serpins from each clade is shown in Figure 1B.

93 Structural studies demonstrated that the RCL region contains a scissile bond between residues  
94 P1 and P1’, that interacts with and can be cleaved by the target protease (Li *et al.* 1999). Upon  
95 cleavage, the reactive center loop of serpin inserts into the  $\beta$ -sheet A. This conformational  
96 change makes serpins thermodynamically more stable (Dementiev *et al.* 2006; Gong *et al.* 2015).  
97 In addition, a fluorescence study demonstrated that the protease in the complex was also  
98 conformationally distorted (Elliott *et al.* 1996). As a result, the target protease is trapped in a  
99 covalent and irreversible complex with the serpin, and thus is inhibited irreversibly (Irving *et al.*  
100 2000; Stein & Chothia 1991). This process is illustrated in Figure 2A, and a structure model  
101 showing covalent serpin-protease complex is shown in Figure 2B.

102 It is worth noting that co-factors are sometimes needed or can enhance serpins’ inhibitory  
103 functions. For example, the glycosaminoglycan heparin, a known anti-coagulant, enhances  
104 inhibition of cathepsin L by serpin B3 and B4 (Higgins *et al.* 2010). It is also interesting to know  
105 that serpins can be secreted or intracellular, thus may also impact their targeted proteases and  
106 ways of functions. For instances, the secreted serpins such as SERPINA1 and SEPRINA3 can  
107 inhibit inflammatory response molecules; while the intracellular serpins such as SERPINB9 acts  
108 on cytosolic proteases thus participate in cellular events (Law *et al.* 2006; Lomas 2005; Sun *et al.*  
109 1996). This does not mean that different forms of serpins have distinct functions, in fact many

110 intracellular serpins participate in inflammatory responses, or vice versa; it is just something we  
111 shall keep in mind when discuss the underlying mechanisms of serpin functions as defense  
112 factors and pathogenic agents.

113

#### 114 **Serpins Serve As Host Defense Factors**

115 The defense strategies of serpins derived from the host are variable, including direct inhibition of  
116 pathogen proteases, inhibition of pathogen binding, and enhancement of host immune cell  
117 functions. Here we discuss mechanisms of how serpins function as host defense factors in a few  
118 representative organisms, such as humans, insects and plants.

119

#### 120 ***Serpins in Vertebrates***

121 In humans and other vertebrates, neutrophil extracellular traps (NETs) are web-like DNA  
122 structures extruded into the extracellular environment by activated neutrophils. NETs are thought  
123 to represent a unique defense strategy against microbial infection. A serpin superfamily member  
124 expressed by macrophages and neutrophils is SerpinB1. It is capable of restricting NET  
125 production. Studies indicate that SerpinB1 inhibits neutrophil elastase, cathepsin G, and  
126 proteinase 3 (Farley *et al.* 2012). More recently, serpins expressed at the mucosal surface have  
127 been linked to inhibition of HIV binding, replication and reduction of inflammation of  
128 susceptible cells. These serpins, together with other protease inhibitors, are found to be expressed  
129 at the epithelial layer of the female genital tract, and thus are considered as essential in the  
130 frontline defense against infection. In addition, their potential applications in disease treatment  
131 have also been explored (Aboud *et al.* 2014).

132 Serpins are also found to regulate defense reactions in other mammalian species, such as  
133 mouse. A serpin superfamily member  $\alpha$ 1-antitrypsin promotes lung defense against  
134 *Pseudomonas aeruginosa* (PA) infection in mice. A study by Jiang *et al.* demonstrated that the  
135 underlying mechanism by which  $\alpha$ 1-antitrypsin reduces lung bacterial infection is through  
136 inhibiting neutrophil elastase-mediated host defense protein degradation (Jiang *et al.* 2013).  
137 Potential therapeutic application of  $\alpha$ 1-antitrypsin to both humans and mice during PA infection  
138 has been proposed.

139

#### 140 ***Serpins in Invertebrates***

141 Serpins have been described in invertebrates, particularly insects. Insects utilize innate immunity  
142 as the major defense system against pathogen invasion. The immune responses include hemocyte  
143 clotting, melanization and antimicrobial peptide expression (Meekins *et al.* 2017). To set off  
144 these innate responses, cascades of serine proteases activation must be triggered, and these  
145 proteolytic cascades are tightly regulated by serpins. Many serpin genes have been identified in  
146 species of insects including 34 in *Bombyx mori*, 32 in *Manduca sexta*, 31 in the beetle *Tribolium*  
147 *castaneum* and 29 in *Drosophila melanogaster*. Insect serpins are found expressed in various  
148 organs including fat bodies, midgut and hemocytes (Meekins *et al.* 2017). The majority of these  
149 insect serpins are believed to be related to innate immunity. For example, when *Bombyx mori*  
150 was challenged by pathogens such as *Micrococcus luteus* and *Serratia marcescens*, the

151 expression of serpin6 (BmSerp6) was increased significantly (Li et al. 2017). It was found that  
152 BmSerp6 directly inhibited the expression of the antimicrobial proteins drosomycin and  
153 gloverin2, and the prophenoloxidase (PPO) activity in the melanization cascade.

154 In studies of *Manduca sexta*, researchers demonstrated that serpin-1 could form a complex  
155 with the serine protease hemolymph protease 8 (HP8), to inhibit the activation of the Toll  
156 pathway (An et al. 2011) during bacterial infection. Serpin-7 was found to inhibit  
157 prophenoloxidase-activating protease-3 (PAP3) in the melanization pathway to down-regulate  
158 innate immune responses (Suwanchaichinda et al. 2013). In addition, more recent studies  
159 showed that *Manduca sexta* serpin-1, 4, 9, 13 and serpin-3, 5, 6 were all able to complex with  
160 pro-hemolymph protease 1 (ProHP1), which is a key proteinase in innate immunity of insects  
161 (He et al. 2017).

162 In *Drosophila*, necrotic protein is one of the many serpins that have been related to innate  
163 immunity. Necrotic protein inhibits the clip domain of serine protease persephone, and  
164 *Drosophila* with necrotic protein mutations constitutively express anti-microbial peptide  
165 drosomycin in the Toll inflammatory signaling pathway (Robertson et al. 2003).

166 Thus, most insect serpins negatively regulate innate immunity by inhibiting serine proteases  
167 that are essential for immune responses. In addition, several studies revealed that insect serpins  
168 could also possess direct anti-pathogen activity upon infection (Levashina et al. 1999). For  
169 example, serpin protein SRPN6 from *Anopheles gambiae* was highly up-regulated in epithelia  
170 immediately after bacterial and parasitic exposures. The AgSRPN6 acts directly on parasite  
171 clearance by inhibiting melanization and promoting parasite lysis (Abraham et al. 2005).

172

### 173 ***Serpins in Plants***

174 Serpins are present in almost all land plants (Roberts et al. 2011). However, the functions of  
175 plant serpins remain to be characterized. *In vitro* studies have demonstrated the protease-  
176 inhibiting activities of plant serpins, but the lack of target chymotrypsin-like proteases within  
177 plants suggests that plant serpins may target digestive proteases from invaded pathogens or  
178 parasites. In addition, there is abundant accumulation of serpins in plant seeds endosperm,  
179 phloem of coleoptiles and leaves (Roberts et al. 2003). These localizations of plant serpins  
180 further imply the defensive roles of serpins in plants against exogenous proteases and pathogens.  
181 Although plant serpins may have distinct characteristics from their insect and animal  
182 counterparts, they have been shown to have a role in the pathways regulating the host immune  
183 responses. For example, Yoo et al. demonstrated a serpin protein, *Cucurbita maxima* phloem  
184 serpin-1 (CmPS), had effective elastase-inhibiting activity. They showed that increased  
185 expression of CmPS-1 within the phloem sap was associated with reduced ability of sap-sucking  
186 insect aphids to survive and reproduce (Yoo et al. 2000).

187 Plants serpins are also found to participate in plant immunity as negative regulators of stress-  
188 induced cell death, or so-called hypersensitive response (HR). For instance, *Arabidopsis thaliana*  
189 serpins AtSRP4 and AtSRP5 negatively regulate stress-induced cell death induced by bacteria  
190 (Bhattacharjee et al. 2017). This kind of cell death or response usually occurs at sites where



191 pathogens attempt to invade. Thus the activities of serpins may have a protective role for plants  
192 when facing pathogen attack.

193 In addition to the above mechanisms of serpins acting on proteases to function as host  
194 defense factors, there are also other ways that serpins participate in host defense. For example,  
195 serpins can induce the expression of host antimicrobial peptides and cytokines (*Kausar et al.*  
196 *2017; Zhao et al. 2014*). Serpins can also directly bind to bacterial pathogens and cause  
197 membrane disruptions (*Malmstrom et al. 2009*). Interestingly, even non-inhibitory serpin can  
198 exert antibacterial activity. For example, non-inhibitory serpin ovalbumin-related protein X  
199 possesses antibacterial activity through heparin-binding ability (*Rehault-Godbert et al. 2013*).  
200 All these findings broaden our understanding of the mechanisms of serpin functions as host  
201 defense factors.

202

### 203 **Pathogen-derived Serpins in Infection and Inflammation**

204 As mentioned above, serpins are present in almost all kingdoms of life including microbes and  
205 other pathogenic organisms. Pathogen-derived serpins may facilitate infection or survival of  
206 pathogens, but the mechanisms remain to be fully elucidated. The data indicate that pathogen-  
207 derived serpins are capable of inhibiting host inflammatory proteins or cells, and abrogating host  
208 cell apoptosis. Studies are on-going to find additional strategies used by pathogen-derived  
209 serpins to facilitate infection. In addition, the potential of pathogen-derived serpins as novel  
210 candidates of clinical therapies or vaccination has drawn great interest from scientists and  
211 physicians.

212

### 213 ***Serpins from Viruses***

214 Viral genomes are usually kept at minimum scales to fit their unique life style. Thus the presence  
215 of viral serpins must be essential for the survival and/or function of the virus. In fact, researchers  
216 have identified several serpins that are required for virulence and infectivity (*Nathaniel et al.*  
217 *2004*). In the myxoma viruses, three serpins, SERP1, SERP2 and SERP3 have been identified  
218 (*MacNeill et al. 2006*). Similarly, there are three serpins in orthopox viruses, designated SPI-1,  
219 SPI-2 and SPI-3 (*Macen et al. 1996*). In addition, the P1 positions of SPI-2-like and SERP2  
220 serpins contain an aspartyl residue, which indicates their potential targets are mammalian  
221 caspases and the serine protease granzyme B (*Turner et al. 1999*). By inhibiting these host  
222 proteases, the virus may be able to restrain apoptosis in host cells thus down-regulating host  
223 immune responses. Other viruses have also been shown to contain one or more serpins, including  
224 swinepox, lumpy skin disease virus, fowlpox, and members of the rhadinovirus genus. Recently,  
225 a baculovirus serpin Hesp018 has been identified in the *Hemileuca* species nuclear polyhedrosis  
226 virus (NPV). This serpin protein has been suggested to abrogate host cell apoptosis, resulting in  
227 accelerated production of virus in Sf9 insect cells (*Ardisson-Araujo et al. 2015*).

228 The ability of viral serpins to abrogate host immune systems has been proposed as a strategy  
229 to treat certain diseases. One such example is the proposed application of the viral serpin Serp-1  
230 to treat acute unstable coronary syndromes (*Lucas et al. 2009*). Several other viral serpins are  
231 being studied for their potential applications as novel anti-inflammatory therapeutics as well



232 (Mangan *et al.* 2017; Zheng *et al.* 2012).

233

### 234 ***Serpins from Bacteria***

235 *Tannerella forsythia* is an anaerobic, gram-negative bacteria species that usually resides in the  
236 human mouth and contributes to chronic periodontitis (Pereira *et al.* 2017). To inhibit host  
237 endopeptidases, *T. forsythia* secretes a serpin-type protease inhibitor called miropin, which  
238 irreversibly inhibits serine and cysteine endopeptidases of the host (Goulas *et al.* 2017).  
239 Phylogenetic analysis of this serpin protein shows that it does not follow a vertical descent model,  
240 indicating miropin may arise from the host by horizontal gene transfer. In fact, the studies of  
241 serpins from human commensal bacteria and their therapeutic applications have become exciting  
242 research areas. For instance, Mkaouar *et al.* characterized two novel serpins from the human  
243 gastrointestinal tract commensal bacteria. These two serpins are called siropin-1 and siropin-2.  
244 These two serpins are found to preferentially inhibit two human serine proteases, neutrophil  
245 elastase and proteinase 3, that are associated with human inflammatory bowel disease (Mkaouar  
246 *et al.* 2016). Thus siropins or other serpins from human commensal bacteria have been suggested  
247 as novel therapeutics against human inflammatory diseases.

248

### 249 ***Serpins from Parasites***

250 Parasites are organisms that live in another organism and may cause major public health  
251 problems such as zoonotic diseases. Usually the parasites need to evade host defense system in  
252 order to survive. Serpins are found to be important during host-parasite interaction, and parasites  
253 utilize their serpins to facilitate infection and survival in the host. For example, cattle tick  
254 *Rhipicephalus microplus* encodes at least 24 serpins, of which RmS-3, RmS-6, and RmS-17 were  
255 identified in the saliva and later confirmed to inhibit pro-inflammatory and pro-coagulatory  
256 proteases of the host (Tirloni *et al.* 2014). In a follow-up study, rRmS-3 was found to inhibit  
257 chymotrypsin, cathepsin G, and pancreatic elastase. Among these serpins, rRmS-6 was found to  
258 inhibit trypsin, chymotrypsin, factor Xa, factor XIa and plasmin; while rRmS-17 inhibited  
259 trypsin, cathepsin G, chymotrypsin, plasmin and factor XIa (Tirloni *et al.* 2016). This study also  
260 claimed that polyclonal antibodies to saliva proteins of *Amblyomma americanum*, *Ixodes*  
261 *scapularis* and *Rhipicephalus sanguineus* were able to cross-react with these three *R. microplus*  
262 saliva serpins. These findings suggest serpins from pathogens could be applied as novel  
263 candidates of vaccination (de la Fuente *et al.* 2007).

264 Researchers have identified several serpins associated with *Trichinella spiralis* including  
265 Tsp03044 and TspAd5 (Knox 2007; Zhang *et al.* 2016). Both of them inhibit trypsin,  $\alpha$ -  
266 chymotrypsin and pepsin of mammals. These data support the inference that serpins from  
267 parasites facilitate invasion into host tissues (Zhang *et al.* 2016).

268 Similarly, *Schistosoma mansoni* has at least eight serpins. Among those, Smpi56 and SmSPI  
269 have been characterized. Smpi56 was purified from extracts of adult *S. mansoni*, and is able to  
270 inhibit neutrophil elastase, pancreatic elastase and an endogenous cercarial protease (Ghendler *et*  
271 *al.* 1994; Quezada & McKerrow 2011). SmSPI, also known as *S. mansoni* serpin isoform 3, was  
272 found to inhibit chymotrypsin, neutrophil elastase and porcine pancreatic elastase (Pakchotanon

273 *et al.* 2016). In addition, SmSPI was found predominantly expressed in the head gland of the  
274 parasite, as well as on its spines. These findings suggest that serpins from *S. mansoni* facilitate  
275 intradermal and intravenous survival of this pathogen.

276

### 277 **Potential Crosstalk between Host and Pathogen Serpins**

278 As we discussed above, under most circumstances serpins target proteases to modulate  
279 inflammatory responses during host-pathogen interactions. Yet it is still of great interest to know  
280 whether serpins function through serpin-serpin interactions. In fact, there is evidence that serpins  
281 can be inactivated through polymerization (*Gettins & Olson 2016*). Serpin polymerization occurs  
282 when the RCL region of one serpin docks into the  $\beta$ -sheet A of another serpin to form an inactive  
283 serpin polymer. Actually, this kind of serpin inactivation is not uncommon and many examples  
284 are found in human deficiency and diseases. For example, the Z allele of SERPINA1  
285 accumulates in patients' liver through serpin polymerization (*Lomas et al. 1992*). Thus, we  
286 would not be surprised to find out in the future that pathogen serpins utilize this mechanism to  
287 inactivate host serpins, or vice versa, during infection.

288

### 289 **Conclusions**

290 Almost all organisms express serpins, and serpins play critical roles in host-pathogen interactions  
291 and regulation of inflammatory responses. The evidence indicates that serpins of the host provide  
292 protection and those of the pathogens enhance infectivity. A brief summary of representative  
293 serpin functions during host-pathogen interactions is shown in Figure 3.

294 On the basis of existing studies, we conclude that serpins, either from hosts or pathogens,  
295 modulate inflammatory responses by inhibiting target proteases associated with host-pathogen  
296 interactions. There is also evidence indicating potential cross interactions between host and  
297 pathogen serpins. In addition to all that, there are a number of studies demonstrating that serpin-  
298 proteases complexes can lead to downstream signals that result in responses such as  
299 inflammation, cytoskeleton rearrangement, proliferation and apoptosis. For example, R1-Anti-  
300 chymotrypsin-cathepsin G complexes have been shown to stimulate production of interleukin 6  
301 and activation of NADPH oxidase (*Kurdowska & Travis 1990; Schuster et al. 1992*). More  
302 recently, serpin  $\alpha$ -1 antitrypsin was found to be able to form complexes independently of the  
303 inhibitory site with neutrophil-expressed pro-inflammatory leukotriene B4. The resulting  
304 complexes modulate downstream signaling events, and augmentation of serpin  $\alpha$ -1 antitrypsin  
305 was suggested as a potential therapy for inflammatory diseases (*O'Dwyer et al. 2015*). These  
306 signaling properties of serpin-protease complexes shed light on the versatility of serpins  
307 modulating inflammatory responses. Another interesting yet poorly explored aspect is the  
308 activities of the peptide derivatives resulting from proteolytic cleavage of the reactive center loop  
309 (RCL) of serpins. A few studies have demonstrated that the peptides derived from serpin RCL  
310 have expanded functions such as anti-inflammatory and antimicrobial activities (*Ambadapadi et*  
311 *al. 2016; Andersson et al. 2004*). Further explorations are needed for additional activities and  
312 potential applications of these peptide derivatives.

313 There is no doubt that further studies of serpins will identify many more biological targets and

314 underlying molecular mechanisms. The study of serpins will remain an important area for basic  
315 research, as well as for clinical applications.

316

317

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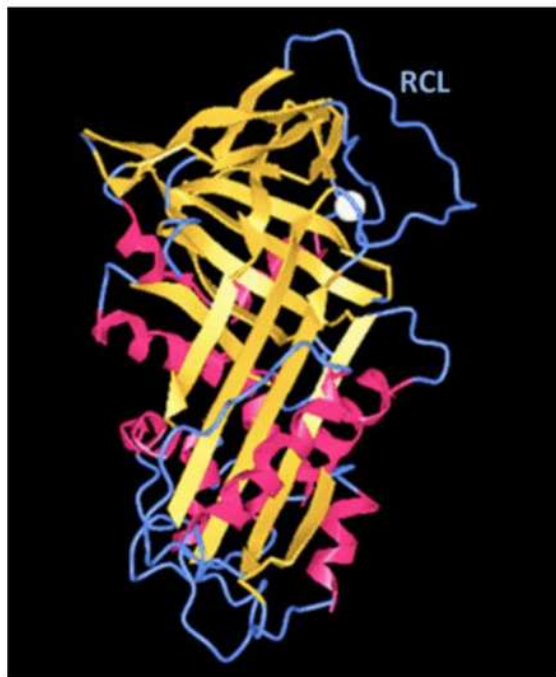
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# Figure 1

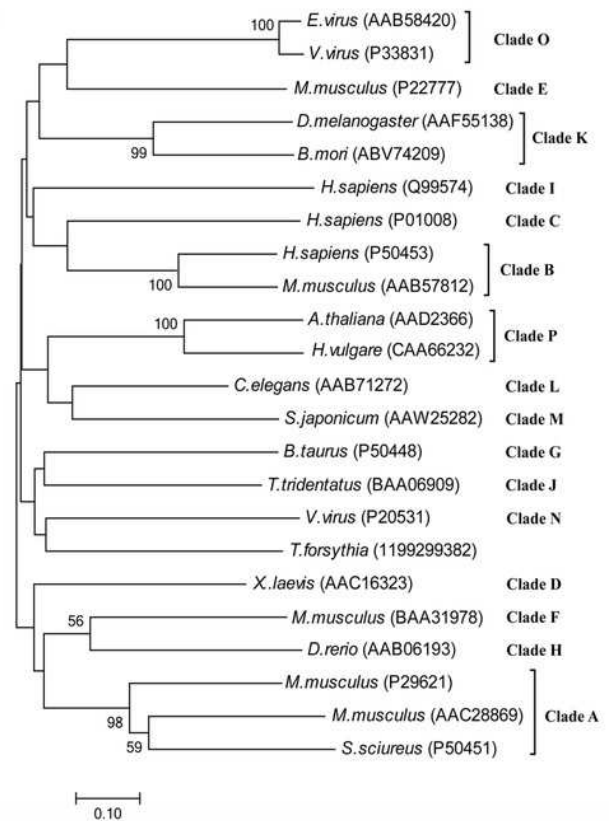
Serpin structure and phylogenetic tree

**Serpin structure and phylogenetic tree. (A)** Structure of the serpin alpha-1 antitrypsin. Human alpha-1 antitrypsin is representative of serpin structures. It contains  $\alpha$ -helices (red),  $\beta$ -sheets (golden) and a reactive center loop (RCL, the upright blue region). (PDB: 3NE4). **(B)** Phylogenetic tree of serpin superfamily. The neighbour-joining tree is based on serpin protein sequences and different clades are represented by a single identifier (e.g., Antithrombin III, P01008), where possible. The phylogenetic analysis was performed using MEGA version 7.0. Analysis was done on 1000 bootstrapped datasets and values of >50% are shown.

(A)



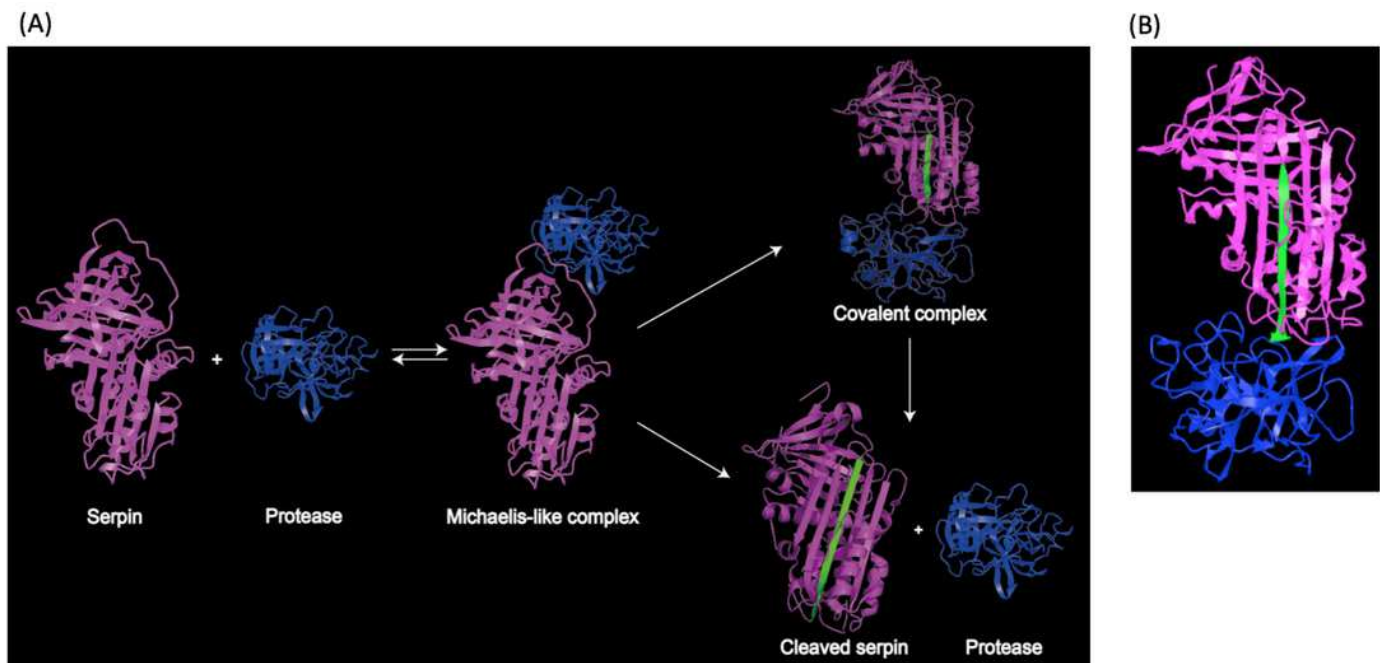
(B)



## Figure 2

### Representation of serpin-protease interaction

Representation of serpin-protease interaction. **(A)** Proposed process of serpin-protease interaction. A serpin (magenta) interacts with targeted protease (blue), and the Michaelis-like complex of serpin and protease is formed. The complex either undergo peptide bond hydrolysis resulting in a kinetically trapped loop-inserted covalent complex (inhibitory pathway), or a cleaved serpin and free protease (non-inhibitory/substrate pathway). The cleaved and inserted RCL is highlighted in green. Serpin-protease complex is stable. Possibility of transition from covalent complex to cleaved form exists yet slim, since complex *in vivo* would be cleared long before complex decay could occur. **(B)** Structure of stable serpin- protease complex (PDB: 2D26). The complex is formed by serpin $\alpha$ 1PI (magenta) and protease elastase (blue). The inserted RCL is highlighted in green.



## Figure 3

### Summary of serpin functions in host-pathogen interactions

Summary of serpin functions in host-pathogen interactions. Hypothesis of protective mechanisms offered by host serpins (on the left, blue), and pathogenic mechanisms exerted by pathogen-derived serpins (on the right, red). Host serpins may act directly or indirectly upon pathogen infections. The representative mechanisms include inhibiting pathogenic digestive proteases, promoting host antimicrobial peptide expression and so on. Pathogen-derived serpins also utilize various mechanisms and representative ones are listed.

