Pathophysiological functions of cathepsin D: Targeting its catalytic activity versus its protein binding activity?
Olivier Masson, Anne-Sophie Bach, Danielle Derocq, Christine Prébois, Valérie Laurent-Matha, Sophie Pattingre, Emmanuelle Liaudet-Coopman

To cite this version:

HAL Id: inserm-00491874
http://www.hal.inserm.fr/inserm-00491874
Submitted on 14 Jun 2010

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Pathophysiological functions of cathepsin D: targeting its catalytic activity versus its protein binding activity?

Olivier Masson, Anne-Sophie Bach, Danielle Derocq, Christine Prébois, Valérie Laurent-Matha, Sophie Pattingre and Emmanuelle Liaudet-Coopman

IRCM, Institut de Recherche en Cancérologie de Montpellier, Montpellier, F-34298, France; INSERM, U896, Montpellier, F-34298, France; Université Montpellier1, Montpellier, F-34298, France; CRLC Val d’Aurelle Paul Lamarque, Montpellier, F-34298, France.
Tel (33) 467 61 24 23; FAX (33) 467 31 37 87; E-mail: e.liaudet@valdorel.fnclcc.fr

Keywords: cathepsin D, protease, cancer, proliferation, apoptosis
Cathepsin D and inhibitors

Abstract
The lysosomal aspartic protease cathepsin D (cath-D) is over-expressed and hyper-secreted by epithelial breast cancer cells. This protease is an independent marker of poor prognosis in breast cancer as it is correlated with the incidence of clinical metastasis. In normal cells, cath-D is localized in intracellular vesicles (lysosomes and endosomes). In cancer cells, overexpressed cath-D accumulates in cells, where it may affect their degradative capacities, and the pro-enzyme is hypersecreted in the tumor micro-environment. In addition, during apoptosis, lysosomal cath-D is released into the cytosol, where it may interact with and/or cleave pro-apoptotic, anti-apoptotic, or nuclear proteins. Several studies have shown that cath-D affects various different steps in tumor progression and metastasis. Cath-D stimulates cancer cell growth in an autocrine manner, and also cath-D plays a crucial paracrine role in the tumor micro-environment by stimulating fibroblast outgrowth and tumor angiogenesis. A mutant D231N-cath-D, which is devoid of catalytic activity, remained mitogenic, indicating an additional action of cath-D by protein-protein interaction. Targeting cath-D in cancer may require the use of inhibitors of its catalytic activity, but also the development of new tools to inhibit its protein binding functions. Thus, elucidation of the mechanism of action of cath-D is crucial if an appropriate strategy is to be developed to target this protease in cancer. The discovery of new physiological substrates of cath-D using proteomic approaches can be expected to generate new critical targets. The aim of this review is to describe the roles of the cath-D protease in cancer progression and metastasis, as well as its function in apoptosis, and to discuss how it can be targeted in cancer by inhibiting its proteolytic activity and/or its binding protein activity.
1. Introduction

Proteases irreversibly hydrolyze the peptide bond in proteins, which results in an important and irreversible post-translational modification. The human genome encodes over 569 proteolytic enzymes or homologs, which constitute the second largest class of human enzymes. Proteases are assigned to five classes on the basis of the active site residue that executes the nucleophilic attack on the target peptide bond: aspartic, cysteine, serine, threonine and metallo-proteinases. These proteases are implicated in normal physiological processes, but deregulation of their expression and/or enzyme activity in disorders such as cancer has profound consequences.

Different families of proteases have been implicated in motility, invasion, extravasation, proliferation and metastasis: the serine proteases (uPA, uPAR and PAI1) [1], the metalloproteinase family [2], the cysteine cathepsins (cathepsin-B and cathepsin-L) [3], and the aspartic cathepsin-D (cath-D) [4], respectively. Cathepsin proteases are lysosomal hydrolases that degrade proteins at acidic pH in the lysosomes, or extracellularly in the matrix. Cathepsins can be divided into three subgroups, based on their active-site amino acid (i.e., cysteine (B, C, H, F, K, L, O, S, V, X, and W), aspartate (D and E), or serine (A and G) cathepsins). The possible involvement of cysteine and aspartic cathepsins in cancer has been the subject of more debate than that of metalloproteases and serine proteases. This might result from the assumption that only secreted proteases that are proteolytically active at neutral pH play an active role in cancer, whereas cathepsins, which require a more acidic pH to be proteolytically active, are thought to be likely to play only a minor role. However, it has been demonstrated that cathepsins are hypersecreted in cancer, and cath-B and cath-D have been described as being associated with the cell surface [5, 6]. Recent studies using transgenic mouse models have stimulated fresh interest in the fundamental roles of cathepsins in cancer [7-10]. Although historically studies have tended to focus on the role of lysosomal proteases
within the endocytic and lysosomal compartments, recent discoveries have shown that these proteases play a critical role in other intracellular compartments, such as the cytosol [11] or the nucleus [12], and within the extracellular milieu in the tumoral stroma [13]. It has become clear that their pattern of expression and their substrate specificities are more complex than was originally envisaged. In cancer, lysosomal proteases are overexpressed and their cellular localizations are profoundly altered, leading to major changes in their targets and consequently in their biological activities.

In addition, cathepsins, metalloproteases and serine proteases act in a cascade-like manner and as part of a proteolytic pathway rather than simply functioning individually [14]. Elucidating the cascade of enzymatic activities that contribute to overall proteolysis during carcinogenesis may identify rate-limiting steps or pathways that could be targeted by anti-cancer treatments [14]. The proteolytic cascade of activation of the different classes of proteases strongly suggests that anti-cancer strategies intended to target several classes of proteases simultaneously might be more promising than those that target a single protease or class of proteases.

Recent studies have focused on extracellular proteases as primary targets for drug discovery, because of their differential expression in many pathophysiological processes, including cancer, cardiovascular conditions, and inflammatory, pulmonary, and periodontal diseases [3, 15]. Interestingly, new extracellular inhibitors of metalloproteases, serine proteases and cysteine proteases are currently under clinical investigation [15]. The aim of this review is to present the role of the cath-D protease in cancer progression and metastasis, as well as its function in apoptosis, and to discuss how it could be targeted in cancer by inhibiting its proteolytic activity and/or its protein binding activity.

2. Structure and trafficking of cath-D
Cath-D [E.C. 3.4.23.5] is a ubiquitous, lysosomal, aspartic endo-proteinase that requires an acidic pH to be proteolytically active (Figure 1). The human cath-D gene contains 9 exons, and is located on chromosome 11p15 [16]. During its transportation to lysosomes, the 52-kDa human pro-cath-D is proteolytically processed to form a 48-kDa, single-chain, intermediate which is an active enzyme located in the endosomes. Further proteolytic processing yields the mature active lysosomal protease, which is composed of both heavy (34 kDa) and light (14 kDa) chains. Cysteine cathepsins are known to be implicated in cath-D processing [17-19]. The involvement of cath-B and L has been shown more recently [19-21]. The human cath-D catalytic site includes two critical aspartic residues (amino acids 33 and 231) located on the 14-kDa and 34-kDa chains, respectively (Figure 1) [22]. Cath-D, like other aspartic proteases such as renin, chymosin, pepsinogen, has a bilobed structure. The crystal structures of the native and pepstatin-inhibited form of mature human cath-D [22-25] have been shown to have a high degree of tertiary structural similarity with other members of the aspartic protease family (e.g. pepsinogen and human immunodeficiency virus protease). The high-resolution structure of pro-cath-D remains to be elucidated. In estrogen receptor (ER)-positive breast cancer cell lines, cath-D is highly up-regulated by estrogens and growth factors (i.e. IGF1, EGF, insulin) [26]. In ER-negative breast cancer cell lines, cath-D is constitutively overexpressed. The mechanism for cath-D overexpression in ER-negative breast cancer cells may involve local reorganization of the chromatin structure of the cath-D promoter [27]. Cath-D overexpression leads to the hypersecretion of the 52-kDa, proteolytically-inactive pro-enzyme, and the accumulation of intracellular cath-D [28]. Release of mature cath-D by exocytosis has been observed in specialized cells [29]. Mannose-6-phosphate (M6P) receptors are involved in cath-D lysosomal routing, and in the cellular uptake of secreted pro-cath-D, although cath-D may also be targeted to the lysosomes, and undergo endocytosis independently of M6P receptors (Figure 1) [30, 31]. The LRP1 receptor has been excluded as
a possible receptor mediating the alternative endocytosis of pro-cath-D on the basis of the inability of the protein chaperone RAP, which competes with ligands that bind to the alpha chain of LRP1, to prevent cath-D endocytosis [32]. It has recently been shown that sortilin functions as an alternative sorting receptor to the M6P receptors for cath-D and cath-H [33].

3. The tight control of cath-D expression and catalytic function is fundamental in normal cells

During fetal development, the level of cath-D increases gradually in all tissues, suggesting a gradual maturation of the lysosomal system [34]. A reduction of cath-D expression or catalytic activity leads to devastating neurodegenerative disorders. Cath-D knockout mice die shortly after birth, and display a neuronal accumulation of ceroid lipofuscin, accompanied by neurodegeneration in the retina and central nervous system, and the accumulation of autophagic vacuoles [35-38]. Cath-D-deficient Drosophila recapitulates the key features of neuronal ceroid lipofuscinoses (NCLs) [39]. Congenital cath-D mutations leading to reduced expression of cath-D and/or the production of enzymatically-inactive protein result in typical NCL in dogs and humans [40-45]. More recently, cath-D deficiency has been shown to be associated with Parkinson disease [46].

In contrast, an increase in cath-D expression can also lead to fatal disorders. A recent study indicates that increased cardiac cath-D expression and activity induces heart failure. This is attributable to a 16-kDa cath-D-cleaved form of prolactin that mediates postpartum cardiomyopathy [47]. Increased cath-D levels have also been observed in the cerebellum of autistic subjects, suggesting that altered activities of cath-D may play an important role in the pathogenesis of autism [48].

4. Function of cath-D in apoptosis
Cysteine and aspartic cathepsins play key roles in tumor cell death via the mediation of apoptosis [4, 11, 49-51]. The function of cath-D in apoptosis needs further investigation, since this protease has both anti-apoptotic and pro-apoptotic functions.

**Anti-apoptotic characteristics of cath-D**

Even though cath-D gene expression outlines the areas of physiological cell death during embryo development [52], cath-D deficiency in mice has revealed its anti-apoptotic function under physiological conditions [35-37]. Indeed, cath-D knock-out mice developed apoptosis in the thymus and in the retina [35-37]. Some other studies have also suggested that cath-D may have an anti-apoptotic role in cancer. Our own immunohistochemical studies have revealed that xenografts of cancer cells overexpressing cath-D displayed less tumor apoptosis than mock-transfected cancer cells [53]. More recently, cath-D has been shown to protect human neuroblastoma cells from doxorubicin-induced cell death [54].

**Pro-apoptotic characteristics of cath-D**

Cath-D is a key mediator of induced-apoptosis, and its proteolytic activity has often been shown to be involved in this event [49, 55-63]. During apoptosis, mature lysosomal cath-D is translocated to the cytosol due to lysosomal membrane permeabilization (LMP) [56-58, 60, 64, 65]. Cytoplasmic cath-D has been shown to cleave Bid to form tBid [66, 67], which triggers the insertion of Bax into the mitochondrial membrane [62, 68], and leads in turn to the mitochondrial release of cytochrome c into the cytosol, and the activation of pro-caspases 9 and 3 [56, 60, 64, 66]. Cath-D is also involved in caspase-independent apoptosis by activating Bax independently of Bid cleavage, and leading in turn to the mitochondrial release of the apoptosis inducing factor (AIF) [68]. More recently, it has been shown that cath-D can also activate pro-caspase 8, initiating neutrophil apoptosis during the resolution of
inflammation [69]. Interestingly, a recent report indicates the presence of mature cath-D in the nucleus during cell death [70], and it has been proposed that nuclear cath-D may mediate the proteolytic activation of endonuclease 23 during cryonecrotic cell death [71]. Since cath-D is one of the lysosomal enzymes that requires a more acidic pH to be proteolytically-active than lysosomal cysteine enzymes, such as cath-B and cath-L, it is open to question whether cytosolic cath-D is able to cleave the substrate(s) implicated in the apoptotic cascade. In some studies, pepstatin A, an inhibitor of the enzyme, partially delayed the apoptosis induced by IFN-gamma and FAS/APO [55], staurosporin [60, 68], TNF-alpha [55, 66, 72], serum deprivation [73], oxidative stress [56, 57, 59], or when pepstatin A was co-micro-injected with cath-D [64]. Other studies indicate that the effect of a mutant cath-D deprived of catalytic activity was indistinguishable from that of the normal enzyme [61, 74]. Furthermore, microinjection of the inactive precursor pro-cath-D into cytosol confirmed that the pro-apoptotic effect of cytosolic cath-D may be also independent of its catalytic activity [75]. In conclusion, cath-D can promote apoptosis by mechanisms that may be dependent on and/or independent of its active site.

5. Roles of cath-D in cancer

*Cath-D is an independent marker of a poor prognosis in breast cancer*

In the 1990s, several independent clinical studies showed that the cath-D level in primary breast cancer cytosols is an independent prognostic parameter correlated with the incidence of clinical metastasis and shorter survival times [76]. A meta-analysis of studies on node-negative breast cancer [77], as well as a complete study of 2810 patients in Rotterdam [78], indicate that high concentrations of cath-D are an effective marker of aggressiveness. Cath-D is now recognized as an independent marker of poor prognosis in breast cancer associated with metastatic risk [79]. In recent years, independent studies have confirmed the prognostic
value of cath-D in breast cancer [80-88]. The main cath-D producing cells appear to be cancer cells and stromal macrophages [89]. Pro-cath-D is also increased in the plasma of patients with metastatic breast cancer [90-92], indicating that some of the pro-cath-D secreted by tumors can be released into the circulation. Interestingly, proteomic studies have recently confirmed the up-regulation of cath-D in many types of cancer [87, 93, 94].

**Cath-D affects multiple steps of cancer progression and metastasis**

Cath-D is overexpressed and hypersecreted in a multitude of cancer types (breast cancer, ovarian cancer, endometrial cancer, cancer of the head and neck, bladder cancer, malignant glioma, melanoma). In cancer cells, overexpressed cath-D accumulates in cells where it may affect their degradative capacities, and the pro-enzyme is hypersecreted in the tumor microenvironment (Figure 2). Cath-D hypersecreted by cancer cells may affect stromal cell behavior and/or degrade components of the extracellular matrix, thus modifying the tumor microenvironment (Figure 2).

Several reports have indicated that cath-D stimulates cancer cell proliferation [95-101], and increases the metastatic potential [96, 100, 102-104]. Cath-D stimulates cancer cell growth in an autocrine manner [97, 98, 105-107]. Various different mechanisms have been proposed to explain the mitogenicity of cath-D. Intracellular cath-D stimulates high density cancer cell growth by inactivating secreted growth inhibitors, such as heat shock 70 protein [99, 108]. Secreted pro-cath-D may act as a mitogen by competing with IGF2 for interaction with the M6P moieties of the M6P/IGF2 receptor, displacing IGF2 from the IGF1 receptor, and resulting in the activation of the mitogenic IGF1 receptor pathway [109, 110]. Many publications have suggested that the interaction of a part of the cath-D pro-fragment (amino acids 27 to 44) with an unknown cell surface receptor is implicated in its mitogenic function [97, 101, 107, 111-113]. Alternatively, it has also been suggested that the catalytic activity of
secreted cathepsin D may be implicated in releasing growth factors, such as FGF2, from the extracellular matrix [114]. Even though, the extracellular pH in tumors is generally more acidic than that in the corresponding normal tissues [115], the question remains as to whether secreted pro-cathepsin D could be activated extra-cellularly in a sufficiently acidic milieu. We have demonstrated that a mutated $^{D231N}$ cathepsin D, which is devoid of proteolytic activity, was still mitogenic for cancer cells both in vitro, in three-dimensional (3D) matrices, and in athymic nude mice [53, 105], suggesting that cathepsin D can also act by protein-protein interaction [116].

Interactions between stromal and epithelial cells are important in cancer progression and metastasis [117-119]. Stromal and tumor cells can exchange numerous tumor-promoting factors, such as growth factors, cytokines, and proteases. The fibroblast is a major cell type of the stromal compartment and, as such, is intimately involved in orchestrating the stromal side of the dialogue in tissue homeostasis. Cathepsin D is localized on the surface of breast fibroblasts [6], and can be taken up by fibroblasts [30, 120, 121]. Cathepsin D has been shown to play a crucial paracrine role in the tumor micro-environment by stimulating fibroblast outgrowth and tumor angiogenesis [53, 122], and possibly by inhibiting anti-tumor responses [123]. More recently, endothelial cells have been shown to secrete pro-cathepsin D via the action of inflammatory cytokines [124]. A mutant version of cathepsin D (D231N) that was devoid of catalytic activity, still proved to be mitogenic for fibroblasts, suggesting a mechanism involving protein-protein interaction [120].

Interestingly, some reports have indicated that the cysteine lysosomal cathepsins, cathepsin L and cathepsin F, which lack a signal peptide, localize in the nucleus [125, 126]. Nuclear cathepsin L proteolytically processes CDP/Cux transcription factor [12, 125] and histone H3 [127, 128], and has important functions in the control of cell transformation [129, 130] as well as in differentiation [127]. It has been shown that translation initiation at downstream AUG sites within cathepsin L mRNA is the first requirement in the chain of events that leads to the presence
of active cath-L in the nucleus [125]. There is, at present, no clear evidence for the presence of cath-D in the nucleus, and the nuclear function of cath-D in cancer is still unknown. However, our preliminary experiments strongly suggest the presence of cath-D in the nucleus of cancer cells. This may be due to translation initiation at downstream AUG sites within the cath-D mRNA (Figure 2). Indeed, it is worth noting that, like that of cath-L [125], the coding sequence of cath-D contains several AUG codons that are located downstream of the first AUG codons. Alternatively, two recent reports have suggested that cytosolic mature cath-D may reach the nucleus in apoptosis (Figure 2) [70]. One important question concerns the ability of cath-D to act as a functional enzyme at the neutral pH of the nucleus. Since enzymatic activity by cath-D is achieved at acidic pH, we can reasonably assume that cath-D might be only weakly active in the nuclear milieu. However, even limited cath-D activity in the nucleus could be compatible with a role in the proteolytic processing of specific nuclear proteins. In contrast, the optimal activity of cathepsins in the acidic environment of the lysosomes is necessary for the terminal degradation of proteins. Thus, the suboptimal pH that prevails in the nucleus should not be taken as an obstacle, but rather as an important element that enables cath-D to play a role in the limited proteolysis of nuclear proteins. Another possibility is that nuclear cath-D may sequester transcription repressors and/or activators, modulating the composition of the complexes implicated in the tight control of transcription. Our preliminary results indicate that cath-D can indeed interact with a nuclear repressor implicated in cancer. Future studies will clarify whether cath-D participates in the regulation of transcription in cancer by cleaving and/or interacting with nuclear proteins, and thus modulates their activity.

6. Targeting cath-D in cancer
Cathepsins have long been known to play an important role in the progression and metastasis of cancer. Cath-D stimulates cancer cell proliferation, fibroblast outgrowth, tumor angiogenesis, and metastasis. In cancer cells, overexpressed cath-D accumulates in cells where it may affect their degradation capacities, and the pro-enzyme is hypersecreted in the tumor micro-environment (Figure 2). Therefore, inhibiting cath-D action requires the development of inhibitors targeting extracellular cath-D, and/or intracellular cath-D located in different parts of the cell (e.g. intracellular vesicles, cytosol, or nucleus).

**Inhibitors of cath-D proteolytic activity**

In recent years, research interest in the development of potent inhibitors of various aspartic peptidases has arisen, fuelled by the growing evidence of their involvement in human diseases [131], such as that of renin in hypertension [132], γ-secretase in Alzheimer's disease [133], plasmepsins in malaria [134], HIV-1 peptidase in acquired immune deficiency syndrome [135], and secreted aspartic peptidases in Candida infections [136]. As opposed to other proteinases (e.g. serine proteases, metalloproteinases or cysteine cathepsins), no mammalian endogenous lysosomal or cytoplasmic cath-D inhibitor is known to exist. When released into the plasma, cath-D is inactivated by its interaction with α2-macroglobulin at a neutral pH, but not at an acidic pH [137, 138]. Since cath-D requires an acidic pH to be proteolytically active, acidic pH may be the physiological regulator of human cath-D activity.

In normal cells, cath-D is only active in acidic intracellular vesicles, and therefore uncontrolled proteolysis is avoided. However, no endogenous cath-D inhibitor is known to exist at acidic pH. It is worth noting that, in cancer, cath-D hypersecreted into the acidic extracellular tumor microenvironment may have a profound effect on matrix remodeling or extracellular factor proteolysis. Most exogenous cath-D inhibitors are synthetic compounds: peptides and polypeptides produced by micro-organisms, plants and lower animals [139, 140].
Organic compounds that esterify the carboxyl group of the Asn33 or Asp231 are synthetic cath-D inhibitors. Studies coupling the complementary methods of combinatorial chemistry and structure-based design, yielded low nanomolar inhibitors of cath-D [141-145]. Cath-D activity is inhibited by structural analogs of synthetic substrates in which an L-amino acid has been replaced by a D-amino acid [139]. The cath-D propeptide segment, which is cleaved off during zymogen activation, has been reported to inhibit pro-cath-D by blocking the active site at neutral pH [24, 25, 146, 147]. At high pH, a stable conformational species of cath-D exists in which the active site is blocked [25]. More recently, a pH-dependent conformational change has been shown to be mediated by electrostatic switches [24]. Peptide fragments derived from the propeptide have been shown to display some inhibitory potency against mature cath-D, suggesting that the development of new classes of pro-peptide-derived inhibitors of cath-D may be promising [147]. Pepstatin A, an inhibitor of aspartic proteases produced by a micro-organism, is the most potent polypeptide inhibitor of cath-D so far identified [148]. This is a hexa-peptide containing the unusual amino acid, statin (Sta, (3S,4S)-4-amino-3-hydroxy-6-methylheptanoic acid), and has the sequence Iva-Val-Val-Sta-Ala-Sta. It was originally isolated from cultures of various species of Actinomycetes due to its ability to inhibit pepsin at picomolar concentrations. It was later found to be a potent inhibitor of nearly all acidic proteases and, as such, has become a valuable research tool. Pepstatin is commonly used to study the role of cath-D in in-vitro systems and in cells. Some studies have seemed to show that pepstatin A administered in vivo induces a significant reduction in the number of metastases, whereas other studies have not confirmed this effect [149]. Inhibition of cath-D by tripeptides containing statin analogs has also been reported [150]. Cath-D polypeptide inhibitors have also been identified in many plants [139], such as tomato leaves [151] and potato tubers [152]. Cath-D inhibitors are also produced by lower animals, such as equistatin from Actinia equina [153, 154] that can also inhibit cysteine cathepsin activity.
Interestingly, it has been shown that deoxyribonucleic acids (DNA fragments) can inhibit cath-D proteolytic activity [155].

**Inhibitors of cath-D binding activity**

Cath-D can also act by protein-protein interaction. Studies of the role of secreted pro-cath-D as a mitogen through its protein binding activity in cancer suggest the involvement of a part of the cath-D profragment (position 27-44) in an interaction with an unknown cell surface receptor [97, 101, 107, 111-113]. Interestingly, an anti-procath-D antibody directed against peptide 27-44 can reverse the growth of human breast tumors in athymic nude mice [111, 112, 156]. Secreted pro-cath-D may also act as a mitogen via its interaction with the M6P moieties of the M6P/IGF-2 receptor, displacing IGF2 from the IGF1 receptor, and leading to the activation the mitogenic IGF1 receptor pathway [109, 110]. We have demonstrated that a mutant $^{D231N}$cath-D that is devoid of proteolytic activity is still mitogenic for cancer cells and fibroblasts both *in vitro* in three dimensional (3D) matrices, and in athymic nude mice [53, 105]. These findings suggest that pro-cath-D may act as an extracellular binding protein by directly or indirectly triggering an as-yet unidentified cell surface receptor. Our unpublished results also indicate that pro-cath-D hypersecreted by cancer cells triggers fibroblast invasive growth in a 3D matrix by interacting with a newly-identified fibroblastic cell surface receptor (submitted). The GST pull-down experiments revealed that this novel cath-D receptor binds the 52-, 34- and 14-kDa cath-D fragments, but only poorly to the 4-kDa cath-D profragment, indicating that the interaction interface spans both 34- and 14-kDa cath-D sub-units (submitted). Taken together, these observations suggest the importance of targeting extracellular pro-cath-D, and open new perspectives for the therapeutic inhibition of protease function in cancer by means other than the use of classical catalytic activity inhibitors. Because of the pleiotrophic action of secreted cath-D as a binding protein, the best strategy
may be to inhibit the extracellular action of pro-cath-D through the use of neutralizing antibodies, rather than by targeting an individual cath-D partner.

Studies in apoptosis also strongly suggest that mature cytosolic cath-D may have an additional role involving protein-protein interaction. So far, no apoptosis-related binding partner of cath-D has yet been identified. The search for cath-D partners using the yeast-two hybrid approach may elucidate the pro-apoptotic function of cath-D independently of its catalytic activity. Our unpublished results using this approach show that cath-D does indeed interact with a pro-apoptotic constituent of the apoptotic pathway. However, it would be premature to envisage blocking the interaction of cath-D using a component of the apoptotic machinery within the cell.

*Cath-D substrates in cancer*

The discovery of new cath-D physiological substrates is likely to generate new critical targets for cancer therapy. To understand the functions of proteases, it is crucial to identify their substrates. Cath-D cleaves preferentially -Phe-Phe-, -Leu-Tyr-, -Tyr-Leu-, and –Phe-Tyr- bonds in peptide chains containing at least five amino acids at an acidic pH [157]. These peptides contain L-amino acids, and also contain hydrophobic amino acid residues at the site cleaved by cath-D. Recently, proteome-derived database-searchable peptide libraries have been developed to identify endoprotease cleavage sites [158]. This approach may be applicable for cath-D. For a long time the main function of cath-D was thought to be to degrade proteins in lysosomes at an acidic pH. In addition to its established role as a major protein-degrading enzyme in lysosomes and phagosomes, it has been shown that cath-D can also activate precursors of biologically-active proteins, such as prolactin and osteopontin in specialized cells [159-163]. Many cath-D substrates have been reported in *vitro*, but no endogenous substrates of cath-D in cancer have yet been clearly identified. In
proteomics, the set of proteins that can be hydrolyzed by a protease is named its substrate degradome or degradomics [164]. A method termed Terminal Amine Isotopic Labeling of Substrates (TAILS) has recently been developed to identify extracellular and membrane protease substrates using iTRAQ labeling and mass spectrometry [165-167]. This powerful proteomic approach, which permitted the discovery of the MMP-2 substrate degradome [168], can also be applied to the identification of cath-D substrates using cells that do or do not express cath-D.

7. Conclusion

Cath-D is a key protease that affects many fundamental functions in cells. The molecular mechanism by which cath-D affects cancer progression remains largely unknown. Furthermore, we still do not have any specific cath-D inhibitors that could be used to target its action in cancer. Since this protease may also act by protein-protein interaction, it will be crucial to identify its partners in order to develop inhibitors to block its protein binding function.

Acknowledgement

Grant sponsors ‘ANR Jeunes chercheurs Jeunes chercheuses’ ANR-05-JCJC-0215-01, and EU FP6; Grant number: LSHC-CT-2007-037665.
References

Cathepsin D and inhibitors


Cathepsin D and inhibitors

Cathepsin D and inhibitors


S. Zhao, E.R. Aviles, Jr., D.G. Fujikawa, Nuclear translocation of mitochondrial cytochrome c, lysosomal cathepsins B and D, and three other death-promoting proteins within the first 60 minutes of generalized seizures, J Neurosci Res (2010 (in press)).

S. Zhao, E.R. Aviles, Jr., D.G. Fujikawa, Nuclear translocation of mitochondrial cytochrome c, lysosomal cathepsins B and D, and three other death-promoting proteins within the first 60 minutes of generalized seizures, J Neurosci Res (2010 (in press)).

S. Zhao, E.R. Aviles, Jr., D.G. Fujikawa, Nuclear translocation of mitochondrial cytochrome c, lysosomal cathepsins B and D, and three other death-promoting proteins within the first 60 minutes of generalized seizures, J Neurosci Res (2010 (in press)).

S. Zhao, E.R. Aviles, Jr., D.G. Fujikawa, Nuclear translocation of mitochondrial cytochrome c, lysosomal cathepsins B and D, and three other death-promoting proteins within the first 60 minutes of generalized seizures, J Neurosci Res (2010 (in press)).

S. Zhao, E.R. Aviles, Jr., D.G. Fujikawa, Nuclear translocation of mitochondrial cytochrome c, lysosomal cathepsins B and D, and three other death-promoting proteins within the first 60 minutes of generalized seizures, J Neurosci Res (2010 (in press)).

S. Zhao, E.R. Aviles, Jr., D.G. Fujikawa, Nuclear translocation of mitochondrial cytochrome c, lysosomal cathepsins B and D, and three other death-promoting proteins within the first 60 minutes of generalized seizures, J Neurosci Res (2010 (in press)).

S. Zhao, E.R. Aviles, Jr., D.G. Fujikawa, Nuclear translocation of mitochondrial cytochrome c, lysosomal cathepsins B and D, and three other death-promoting proteins within the first 60 minutes of generalized seizures, J Neurosci Res (2010 (in press)).

S. Zhao, E.R. Aviles, Jr., D.G. Fujikawa, Nuclear translocation of mitochondrial cytochrome c, lysosomal cathepsins B and D, and three other death-promoting proteins within the first 60 minutes of generalized seizures, J Neurosci Res (2010 (in press)).


E. Liaudet, M. Garcia, H. Rochefort, Cathepsin D maturation and its stimulatory effect on metastasis are prevented by addition of KDEL retention signal, Oncogene 9 (1994) 1145-1154.


Cathepsin D and inhibitors

Cathepsin D and inhibitors

Cathepsin D and inhibitors

**Figure legends**

**Figure 1. Schematic representation of the human cath-D 52 kDa pro-cath-D sequence**

The locations of the 4-kDa cath-D pro-fragment, 14-kDa light, and of the 34-kDa heavy mature chains are indicated. The intermediate 48-kDa form (not shown) corresponds to non-cleaved 14 + 34 kDa chains. According to [169], 1 corresponds to the first amino acid in mature cath-D. The positions of the 2 aspartic acids of the catalytic site are shown, as are the 2 glycosylated chains carrying M6P motifs that are recognized by the M6P receptors. K, kilodalton.

**Figure 2. Localization of cath-D in cancer cells**

In cancer cells, overexpressed cath-D accumulates in cells where it may affect their degradative capacities. The pro-enzyme is also hypersecreted in the tumor micro-environment. Cath-D hypersecreted by cancer cells may be captured back by both cancer and stromal cells, thus affecting the tumor micro-environment (modulation of stromal cell behavior and/or of components of the extracellular matrix). After lysosomal membrane permeabilization (LMP) during apoptosis, lysosomal mature 34-kDa cath-D released into the cytosol may interact with and/or degrade pro-apoptotic or anti-apoptotic proteins. A cytoplasmin form of cath-D may also be involved in the regulation of transcription in cancer by interacting with nuclear proteins and modulating their activity. K, kilodalton.