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**HAL Id: hal-02094546**

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# Exploiting the S4–S5 Specificity of Human Neutrophil Proteinase 3 to Improve the Potency of Peptidyl Di(chlorophenyl)-phosphonate Ester Inhibitors: A Kinetic and Molecular Modeling Analysis

Carla Guarino,<sup>†</sup> Natalia Gruba,<sup>‡</sup> Renata Grzywa,<sup>§</sup> Edyta Dyguda-Kazimierowicz,<sup>||</sup> Yveline Hamon,<sup>†</sup> Monika Legowska,<sup>‡</sup> Marcin Skoreński,<sup>§</sup> Sandrine Dallet-Choisy,<sup>†</sup> Sylvain Marchand-Adam,<sup>†</sup> Christine Kellenberger,<sup>⊥</sup> Dieter E. Jenne,<sup>#</sup> Marcin Sienczyk,<sup>§</sup> Adam Lesner,<sup>‡</sup> Francis Gauthier,<sup>†</sup> and Brice Korkmaz<sup>\*,†</sup>

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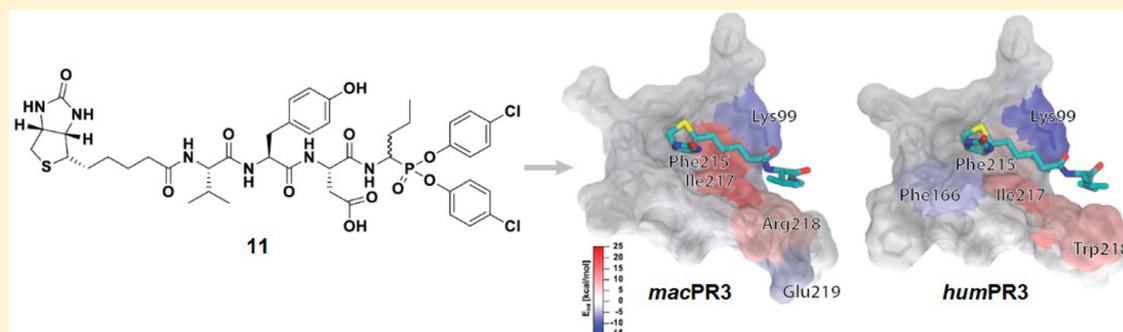
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**S** Supporting Information

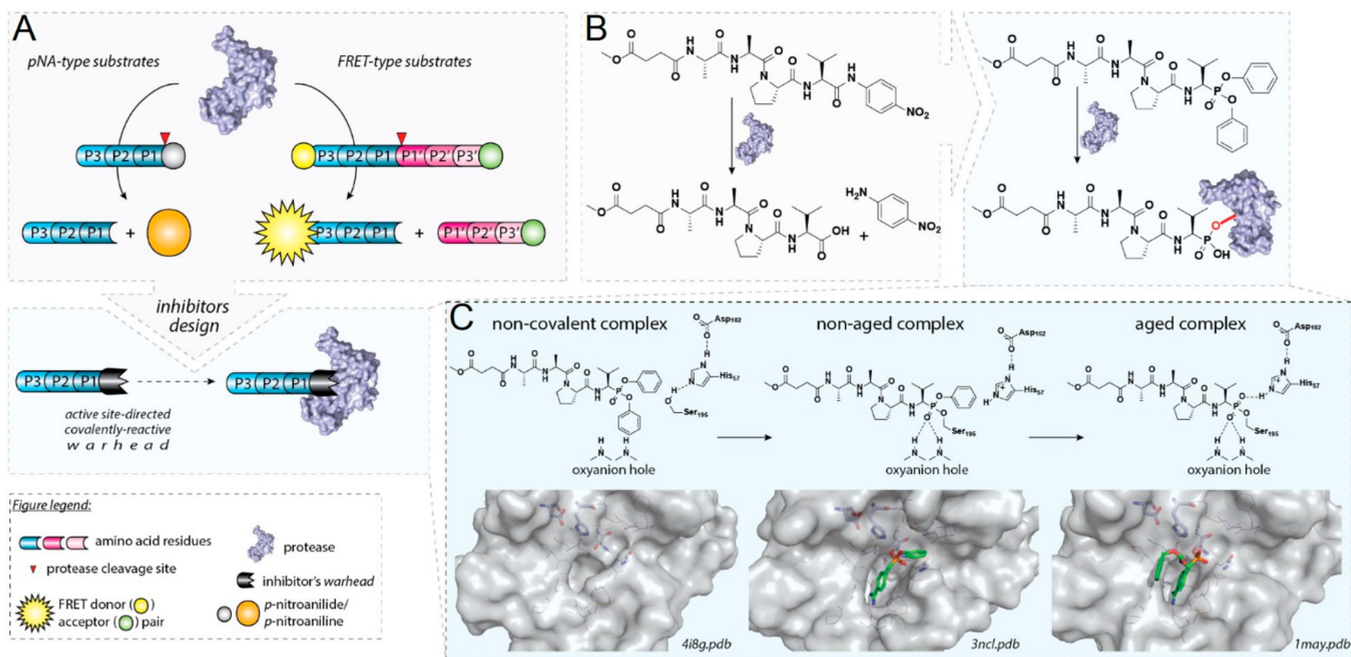


**ABSTRACT:** The neutrophilic serine protease proteinase 3 (PR3) is involved in inflammation and immune response and thus appears as a therapeutic target for a variety of infectious and inflammatory diseases. Here we combined kinetic and molecular docking studies to increase the potency of peptidyl-diphenyl phosphonate PR3 inhibitors. Occupancy of the S1 subsite of PR3 by a nVal residue and of the S4–S5 subsites by a biotinylated Val residue as obtained in biotin-VYDnV<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> enhanced the second-order inhibition constant  $k_{\text{obs}}/[I]$  toward PR3 by more than 10 times ( $k_{\text{obs}}/[I] = 73000 \pm 5000 \text{ M}^{-1} \text{ s}^{-1}$ ) as compared to the best phosphonate PR3 inhibitor previously reported. This inhibitor shows no significant inhibitory activity toward human neutrophil elastase and resists proteolytic degradation in sputa from cystic fibrosis patients. It also inhibits macaque PR3 but not the PR3 from rodents and can thus be used for in vivo assays in a primate model of inflammation.

## INTRODUCTION

Polymorphonuclear neutrophil phagocytes are characterized by the presence of abundant intracytoplasmic granules rich in antimicrobial peptides and proteins involved in innate immunity.<sup>1,2</sup> Azurophilic granules also store four neutrophil serine proteases (NSPs): proteinase 3 (PR3), elastase (NE), cathepsin G (CG), and neutrophil serine protease 4 (NSP-4), which are released into the environment in response to inflammatory stimuli.<sup>1,3</sup> An excess of proteases may be released, however, during chronic inflammation which disrupts the protease–protease inhibitor balance and accelerates proteolysis

of the extracellular matrix.<sup>4,5</sup> The administration of exogenous inhibitors targeting these proteases may thus be an excellent therapeutic strategy to fight inflammation.<sup>5,6</sup> Although the total amount of PR3 in neutrophils is similar to that of NE or CG, its activity is by far less controlled by endogenous inhibitors.<sup>7</sup> Indeed, there is no specific endogenous inhibitor of human PR3 (*humPR3*) and one of its more potent inhibitors,  $\alpha$ -1-proteinase inhibitor ( $\alpha$ 1PI), interacts about 100 times less



**Figure 1.** Design, structure, and mechanism of action of 1-aminoalkylphosphonate diaryl ester inhibitors. (A) General strategy of a substrate-based approach for covalent inhibitors development. (B) Development of a peptidyl-phosphonate inhibitor. (C) Mechanism of serine proteases inhibition by 1-aminoalkylphosphonate diaryl esters together with crystal structures of bovine trypsin (Protein Data Bank (PDB) 4I8G and 1MAY) and human matriptase (PDB 3NCL) at different stages of aging process.

45 rapidly with *humPR3* than with *humNE*.<sup>7</sup> Further, the  
 46 pathophysiological role of *humPR3* is less well understood  
 47 than that of the related *humNE* and CG. Its function as  
 48 autoantigen in granulomatosis with polyangiitis<sup>8–10</sup> and its  
 49 likely involvement in neutrophil apoptosis<sup>11</sup> makes it different  
 50 from its closest homologue *humNE*.

51 *humPR3* closely resembles *humNE* structurally and func-  
 52 tionally with a highly conserved catalytic triad (His57, Asp102,  
 53 and Ser195 residues (*chymotrypsinogen numbering*)) located  
 54 between two similar domains each comprising a six-stranded  $\beta$ -  
 55 barrel.<sup>12</sup> Its pI, however, is somewhat less basic than that of  
 56 *humNE*.<sup>5,13</sup> Several residues on the loops surrounding the  
 57 protease active site assist the catalytic process. Most  
 58 importantly, the backbone amide hydrogens of Gly193 and  
 59 Ser195 that define the oxyanion hole and are located near the  
 60 carbonyl group of the substrate's scissile bond, stabilizing the  
 61 developing partial charge on the tetrahedral intermediate  
 62 during catalysis.<sup>14</sup>

63 The structural analysis of the active site of *humPR3* and  
 64 *humNE* showed that the distribution of charged residues close  
 65 to the substrate binding site (99 loop, 60 loop, 37 loop, and  
 66 autolysis loop) of these two proteases differs notably.<sup>15</sup> Thus,  
 67 *humPR3* contains three charged residues Lys99, Asp61, and  
 68 Arg143 in the active site region.<sup>12</sup> The S1 binding pocket of  
 69 *humPR3* and *humNE* is hemispherical, therefore, both  
 70 preferentially accommodate small hydrophobic residues at the  
 71 P1 position (according to the nomenclature of Schechter and  
 72 Berger (Schechter and Berger, 1967)).<sup>7,13</sup> The S2 subsite of  
 73 *humPR3* differs from that of *humNE* by the presence of a  
 74 solvent accessible Lys at position 99, favoring accommodation  
 75 of negatively charged or polar P2 residues in the deep S2  
 76 subsite of PR3.<sup>12</sup> The Leu99 residue in *humNE* makes the S2  
 77 pocket more hydrophobic. The Lys99 of *humPR3* is conserved  
 78 in the PR3 of higher primates and many artiodactyls but not in  
 79 PR3 of New World monkeys and rodents, whereas the Leu99

of *humNE* is highly conserved in many other species.<sup>7</sup> This  
 80 makes the PR3 specificity of these latter species different from  
 81 that of *humPR3* and explains that rodents are not an  
 82 appropriate animal model for studies related to the biological  
 83 activity of *humPR3*. Another critical residue that makes the  
 84 specificities of *humPR3* and *humNE* different is that at position  
 85 217 in the vicinity of the S4 subsite, where an Ile in *humPR3* is  
 86 replaced by an Arg in *humNE*.<sup>16,17</sup>

87 We have designed and synthesized selective peptidyl-  
 88 diphenyl phosphonate inhibitors based on these structural  
 89 differences between *humPR3* and *humNE* using the sequence of  
 90 an optimized peptide substrate of PR3.<sup>18</sup> Phosphonate  
 91 inhibitors are peptide-based transition state irreversible  
 92 inhibitors which form transition-state-resembling complexes  
 93 with serine proteases.<sup>19–21</sup> The inhibition is initiated by the  
 94 formation of a noncovalent enzyme–inhibitor complex, which  
 95 upon the nucleophilic attack of the Ser195 on the phosphorus  
 96 atom loses one aryloxy group, forming an initial, irreversible  
 97 covalent complex (Figure 1). Further aging followed by  
 98 hydrolysis of a second ester group leads to the formation of  
 99 an aged covalent protease–inhibitor complex stabilized by the  
 100 oxyanion hole.<sup>19</sup> Phosphonate inhibitors are chemically stable  
 101 inhibitors that block selectively serine proteases at low  
 102 concentration under acidic or neutral conditions.<sup>22</sup> Phospho-  
 103 nate inhibitors were designed and developed by anchoring of  
 104 the serine trap to the recognition sequence derived from a  
 105 peptidyl substrate of the target protease (Figure 1). These  
 106 inhibitors which interact covalently with the Ser195 of the  
 107 catalytic triad can also be used as activity-based probes (ABP)<sup>23</sup>  
 108 to visualize membrane-bound or intracellular, proteolytically  
 109 active, serine proteases.<sup>24</sup> Several peptidyl-diphenyl phospho-  
 110 nate inhibitors of *humPR3* have been developed but all were  
 111 more potent toward *humNE*<sup>25,26</sup> until we synthesized the first  
 112 selective chlorodiphenyl phosphonate *humPR3* inhibitors, the  
 113

**Table 1. Rates of Inhibition of *humPR3* and *humNE* by Peptide Phosphonates**

compd	peptide phosphonate esters	[I] $\mu\text{M}$	proteases	
			$k_{\text{obs}}/[I]$ ( $\text{M}^{-1} \text{s}^{-1}$ ) <sup>a</sup>	
			<i>humPR3</i>	<i>humNE</i>
1	Ac-Pro-Tyr-Asp-AlaP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	2	154 ± 3 <sup>b</sup>	ns
2	Bt-Pro-Tyr-Asp-AlaP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	0.06	4168 ± 553 <sup>#</sup>	ns
3	Bt-[PEG] <sub>2</sub> -Pro-Tyr-Asp-AlaP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	0.6	274 ± 12	ns
4	Bt-Val-Tyr-Asp-AlaP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	0.025	17396 ± 835	ns
5	Bt-Leu-Tyr-Asp-AlaP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	0.15	4371 ± 652	ns
6	Bt-Ile-Tyr-Asp-AlaP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	0.15	8698 ± 658	ns
7	Bt-nLeu-Tyr-Asp-AlaP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	0.025	10361 ± 766	ns
8	Bt-nLeu(O-Bzl)-Tyr-Asp-AlaP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	0.1	1744 ± 164	ns
9	Bt-Pro-Tyr-Asp-AbuP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	0.1	4675 ± 438	ns
10	Bt-Pro-Tyr-Asp-nValP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	0.025	18642 ± 705	ns
11	Bt-Val-Tyr-Asp-nValP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	0.01	73258 ± 5342	ns

<sup>a</sup>Values are the means ± SD of three experiments; <sup>b</sup>Values were taken from ref 18. Definition of abbreviation: ns, not significant

114 N-biotinylation of which allows using them ABP to visualize  
115 active *humPR3* in biological samples.<sup>18</sup>

116 Application of PR3 inhibitors as therapeutic tools requires  
117 that they easily reach and interact with their target protease  
118 with great specificity, they resist degradation during their  
119 administration and in situ, and their half-life in the organism is  
120 significant. Using inhibitors as therapeutic tools also requires  
121 that a relevant animal model is available for preclinical studies.  
122 In this work, we first designed and developed new biotin-  
123 peptidyl<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> inhibitors with improved potency of  
124 action toward *humPR3* to use them as versatile pharmacological  
125 tools for assessing protease function in vivo. We focused on  
126 improving the rate constant for inactivation ( $k_{\text{obs}}/[I]$ ) by  
127 molecular docking trials and on analyzing structure–activity  
128 relationships (SAR) to optimize efficacy at a very low dose and  
129 thus make the resulting compound effective for a pharmaco-  
130 logical application. Because PR3 from rodents retain a substrate  
131 specificity that differs from that of human,<sup>27</sup> we then looked for  
132 a relevant in vivo model of inflammation and tested  
133 phosphonate inhibitors on the PR3 from *Macaca fascicularis*.

## 134 ■ RESULTS

135 **Stabilizing Properties of a Biotinylated N-Terminal P4**  
136 **Residue in PR3 Substrates and Inhibitors.** Replacing the  
137 N-terminal acetyl group by biotin (Bt) in Ac-PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-  
138 4-Cl)<sub>2</sub> (**1**) to give Bt-PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (**2**) significantly  
139 improved the  $k_{\text{obs}}/[I]$  value<sup>18</sup> (4168  $\text{M}^{-1} \text{s}^{-1}$  vs 154  $\text{M}^{-1} \text{s}^{-1}$ )  
140 (**Table 1**) and significantly improved the  $K_i$  value of the initial  
141 noncovalent complex (21 vs 3600 nM) (**Table 2**). Accordingly,  
142 the substitution of the N-terminal acetyl group by a biotin in  
143 the paranitroanilide (pNA) substrate Ac-PYDA-pNA increased  
144 the specificity constant  $k_{\text{cat}}/K_m$  by ~6-fold (**Table 3**). We  
145 employed a computational docking approach to explain how  
146 biotin could modulate the interaction between Bt-PYDA<sup>P</sup>(O-  
147 C<sub>6</sub>H<sub>4</sub>-Cl)<sub>2</sub> and the active site of PR3 (**Figure 2A,B**). The lowest  
148 energy binding mode obtained in the docking studies of **2** with  
149 *humPR3* revealed that the biotin moiety is located in the S5  
150 pocket limited by the Lys99, Phe166, Cys168, Arg177, and  
151 Ile217 residues (**Figure 2B**). The entrance into this pocket is  
152 guarded by the Lys99 side chain with its  $\epsilon$ -amino group,  
153 creating a hydrogen bonding with the carbonyl oxygen of the  
154 Bt-Pro4 amide bond. This interaction would facilitate the  
155 correct orientation of both Pro4 and biotin in the S4 and S5  
156 binding sites, respectively. The arrangement of Phe166,

**Table 2. Rates of Inhibition of *humPR3* by Peptide Phosphonates**

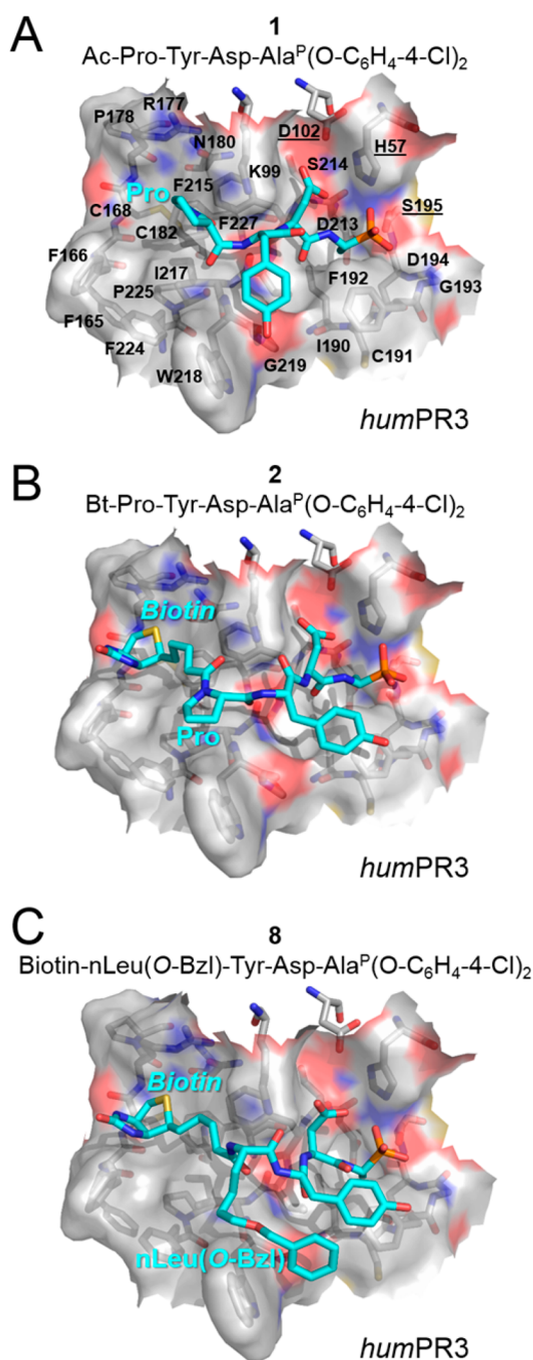
$\text{E} + \text{I} \xrightleftharpoons{K_i} \text{EI} \xrightleftharpoons[k_2]{k_1} \text{E-I}$	
Initial non-covalent complex	Final covalent complex
<b>1</b>	
Ac-Pro-Tyr-Asp-AlaP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	
$K_i$ (nM)	3600 ± 425
$k_2$ (min <sup>-1</sup> )	0.08 ± 0.01
<b>2</b>	
Bt-Pro-Tyr-Asp-AlaP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	
$K_i$ (nM)	21 ± 4.2
$k_2$ (min <sup>-1</sup> )	0.035 ± 0.01
<b>11</b>	
Bt-Val-Tyr-Asp-nValP(O-C <sub>6</sub> H <sub>4</sub> -4-Cl) <sub>2</sub>	
$K_i$ (nM)	5.4 ± 0.14
$k_2$ (min <sup>-1</sup> )	0.15 ± 0.01

**Table 3. Kinetics of Synthetic Substrate Cleavage by *humPR3* and *humNE***

pNA substrates	[S] mM	proteases	
		$k_{\text{cat}}/K_m$ ( $\text{M}^{-1} \text{s}^{-1}$ ) <sup>a</sup>	
		<i>humPR3</i>	<i>humNE</i>
Ac-Pro-Tyr-Asp-Ala-pNA	1	4201 ± 29.7 <sup>b</sup>	nh
Bt-Pro-Tyr-Asp-Ala-pNA	1	25080 ± 141.4	nh
Bt-Val-Tyr-Asp-Ala-pNA	1	34965 ± 99	nh
Bt-Val-Tyr-Asp-nVal-pNA	1	80510 ± 2973	nh

<sup>a</sup>Values are means ± SD of three experiments. <sup>b</sup>Value was taken from ref 18. Definition of abbreviation: nh, not hydrolyzed

Cys168, and Arg177 residues in the S5 subsite creates the 157  
cavity that accommodates the biotin heterocyclic rings (**Figure** 158  
**2B**). The stabilizing role of biotin was confirmed by introducing 159  
a polyethylene glycol [PEG]<sub>2</sub> spacer between the P4 residue 160  
Pro and biotin (Bt-[PEG]<sub>2</sub>-PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (**3**), which 161  
resulted in a dramatic fall of the  $k_{\text{obs}}/[I]$  (**Table 1**). The docking 162  
model shows that the length of the biotin moiety is optimal for 163  
the binding in the S5 pocket, and any spacer between the Pro4 164  
and biotin would not improve the interaction. A biotin at P5 165



**Figure 2.** Proposed putative model of **1** (A), **2** (B), and **8** (C) binding to the active site of *humPR3*. The solvent-accessible surface area of the active site in *humPR3* (PDB 1FUJ<sup>12</sup>) was made transparent to allow the visualization of the residues in stick representation. The single-letter code of residues in the vicinity of the active site is indicated in black. Residues are labeled following the numbering of chymotrypsin. The residues of the catalytic triad H57, D102, and S195 are underlined. The carbon atoms of PR3 and the compounds are shown in white and cyan, respectively. The oxygen, nitrogen, sulfur, and phosphorus atoms are colored in red, blue, yellow, and orange, respectively.

166 was thus retained for the construction of new inhibitors with a  
167 modified peptidyl sequence.

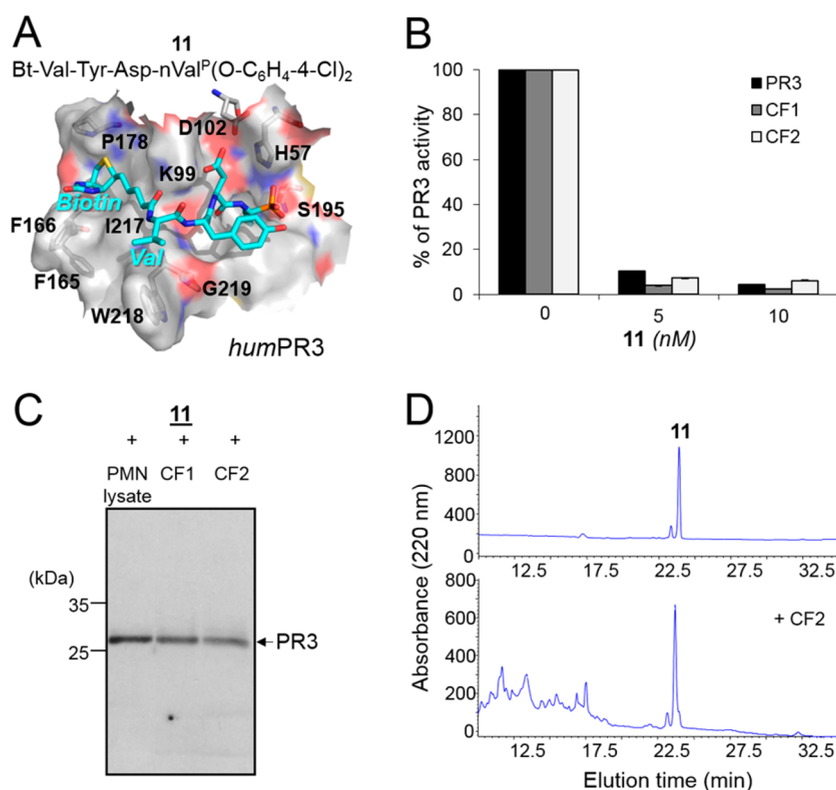
168 **Influence of the P4 Residue on the Inhibitory Activity**  
169 **of Bt-Peptidyl<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> Phosphonate Inhibitors.**  
170 The computational docking study showed that the P4 residue

Pro in **2** was close to solvent accessible hydrophobic Trp218 in  
PR3 (**Figure 2B**). We replaced the P4 Pro by Val (**4**), Leu (**5**),  
Ile (**6**), and norleucine (nLeu) (**7**) to tentatively optimize the  
interaction with the PR3 hydrophobic patch build by residues  
Phe166, Ile217, Phe224, and possibly with Trp218. While Leu  
or Ile at P4 position decreased the inhibitory activity toward  
PR3, the presence of nLeu or Val improved the inhibitory  
activity by  $\sim 2$  and  $\sim 4$  times, respectively (**Table 1**).  
Accordingly, the specificity constant  $k_{\text{cat}}/K_m$  of the pNA  
substrate Bt-VYDA-pNA was also improved (**Table 3**).  
Whatever the substitution at P4 in phosphonate inhibitors  
was, the resulting compound retained no significant inhibitory  
activity toward *humNE* although this protease also prefers a  
hydrophobic residue at this position. Because the S4 subsite is  
composed mainly by side chains of Trp218 and Ile217 and the  
distinctly hydrophobic area (Phe166, Phe224) span beyond this  
position, we decided to probe the existence of interactions by  
substitution nLeu by nLeu(O-Bzl) at P4 in **7** (**8**). However, this  
resulted in more than 10 times lower  $k_{\text{obs}}/[I]$  value (**Table 1**).  
Moreover, the molecular docking model did not confirm the  
interaction between nLeu(O-Bzl) and, as mentioned above, a  
distant hydrophobic area. In fact, the P4 side chain of **8** makes  
contact mainly with Trp218 (**Figure 2C**). The comparison with  
the **2** model (**Figure 2B**) indicates that the introduction of  
more sizable nLeu(O-Bzl) group at P4 does not alter the overall  
mode of binding but affects the placement of inhibitor  
backbone at the S4 subsite, thus preventing hydrogen bond  
formation between Lys99 and the carbonyl oxygen of the Bt-  
nLeu(O-Bzl) amide (**Figure 1C**).

**Influence of the P1 and P4 Residues on the Efficacy of**  
**Bt-Peptidyl<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> Phosphonate Inhibitors.** Un-  
like the S2 subsite of PR3 that preferentially accommodates  
negatively charged P2 residues and is thus essential to confer  
PR3 selectivity,<sup>15</sup> the S1 subsite in PR3 may accommodate a  
variety of residues including norvaline (nVal) and aminobutyric  
acid (Abu) among the favorites. We substituted the P1 alanyl  
residue in the parent inhibitor (**2**) by Abu and nVal. Bt-  
PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (**2**) and Bt-PYDAbu<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>  
(**9**) showed similar efficacy toward PR3 (**Table 1**). However,  
Bt-PYDnV<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (**10**) was  $\sim 4.5$  times more potent  
than **2**.

As expected, the substitution of Pro by Val at P4 in **10** (**11**)  
significantly improved the  $k_{\text{obs}}/[I]$  value, providing the best  
inhibitor of the series with a  $k_{\text{obs}}/[I] = 73000 \pm 5000 \text{ M}^{-1} \text{ s}^{-1}$ .  
This 20-fold increase as compared to **1** resulted from a decrease  
in the  $K_i$  value of the initial equilibrium between PR3 and Bt-  
VYDnV<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (**11**) and an increase of the first-order  
rate constant  $k_2$  producing the final covalent complex (**Table**  
**2**). Combining Val at P4 and nVal at P1 in the pNA substrate  
Bt-VYDnV-pNA also significantly increased the specificity  
constant toward PR3 (**Table 3**).

The computational docking approach employed to examine  
the interaction between Bt-VYDnV<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> and  
*humPR3* (**Figure 3A**) revealed that for the lowest energy  
pose the overall mode of enzyme–inhibitor binding resembles  
the one obtained for Bt-PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (**2**). The biotin  
aliphatic chain interacts with the hydrophobic surface of S5  
subsite, while the biotin rings extend into the terminal cavity of  
this subsite in the manner observed with Bt-PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-  
Cl)<sub>2</sub>. For both models, the P4 residue of inhibitor is located at  
the narrow subsite with the Trp218 on one side and the Lys99  
on the other. Therefore, increased inhibitory potency observed  
for derivatives with Val instead of Pro at P4 may be due to an



**Figure 3.** Inhibition of *humPR3* and stability of **11** in the cell free supernatants of sputa from CF patients. (A) Proposed putative model of **11** in *humPR3* active center. (B) Inhibition of PR3 in a representative CF sputum supernatants. The volume of CF sputum was adjusted to give final concentrations of PR3, 1 nM. Diluted samples were incubated with **11** (5 and 10 nM final) for 20 min at 37 °C. Inhibition was monitored by measuring the residual activity of PR3 on ABZ-VADnVADYQ-EDDnp (20 μM). Purified PR3 (1 nM) was used as control. (C) Selective labeling of PR3 activity in CF sputum and neutrophil lysate. Samples (10 μg of total protein) were incubated for 20 min at 37 °C with **11** (50 nM), and the mixtures were analyzed by WB using extravidin-peroxidase. (D) HPLC profile of the **11** after a 120 min incubation time with samples showing the stability of the inhibitor in sputa from cystic fibrosis patients. Similar results were found in three independent experiments.

234 improved flexibility of this region upon enzyme–inhibitor  
 235 binding. The ε-amino group of Lys99 forms hydrogen bonds  
 236 with Bt-Val4 amide carbonyl oxygen as well as Asp2 carboxyl  
 237 group. Additionally, quantum chemical calculations of inter-  
 238 action energy revealed that Lys99 residue contributes the most  
 239 to binding of Bt-Val4 portion of the inhibitor, as the value of  
 240 the interaction energy due to the presence of this particular  
 241 residue amounts to −16.6 kcal/mol (Table 4). Attracting  
 242 interactions between Bt-Val4 tail of **11** and PR3 residues were  
 243 also found for Phe166 and Val216 (−3.8 and −1.6 kcal/mol,  
 244 respectively). Except for Ile217, Trp218, and Phe215 residues  
 245 that appear to exert unfavorable influence in terms of Bt-Val4  
 246 binding, the remaining PR3 residues promote Bt-Val4 binding  
 247 with the interaction energy not exceeding −1 kcal/mol. It  
 248 should be pointed out that excessively repulsive interactions  
 249 associated with some residues probably arise from the lack of  
 250 quantum chemical refinement of the binding poses obtained  
 251 from docking simulations, as empirical force field based  
 252 methods often employed throughout the docking procedures  
 253 tend to introduce shortened intermolecular contacts.<sup>28</sup> The  
 254 location of the biotin rings into the S5 binding site prevents  
 255 recognition of all these compounds by extravidin by Western  
 256 blotting (WB) under nondenaturing/reducing conditions  
 257 (not shown).

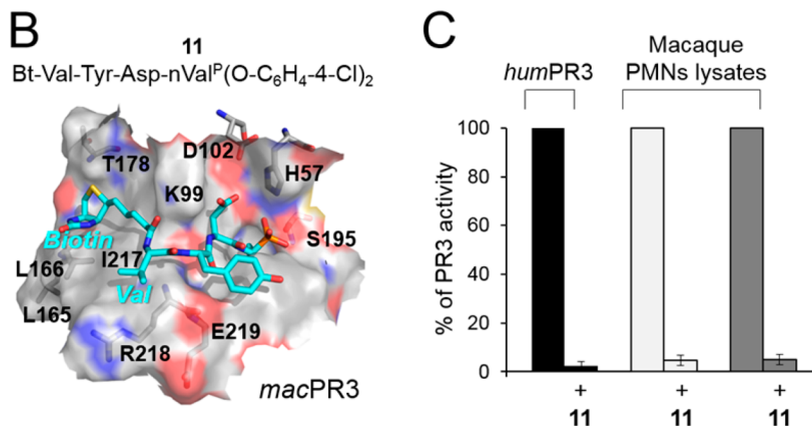
258 **Stability of Bt-VYDnV<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (**11**) in a Bio-**  
 259 **logical Environment.** We then tested the properties of **11** in  
 260 sputa from patients with cystic fibrosis (CF) and measured  
 261 *humPR3* activities of sputum samples before and after

**Table 4. MP2/6-31+G(d) Interaction Energy<sup>a</sup> between Amino Acid Residues Representing *humPR3* or *macPR3* Binding Site and Bt-Val4 Fragment of **11****

<i>humPR3</i> residues	substituted <i>macPR3</i> residues	binding energy	
		<i>humPR3</i>	<i>macPR3</i>
Lys99		−16.6	−10.0
Phe165	Leu	−0.3	−0.4
Phe166	Leu	−3.8	−0.8
Cys168-Cys182		−0.8	−1.2
Asn98-Arg177-Asn180		−0.8	0.3
Pro178	Thr	0.1	0.2
Phe192		−0.1	0.0
Phe215		5.5	24.2
Val216		−1.6	−2.6
Ile217		12.3	15.4
Trp218	Arg	7.6	6.4
Gly219	Glu	−0.3	−3.3
Phe224		−0.6	−0.9
Pro225		−0.6	−0.7
Phe227		−0.8	−1.6

<sup>a</sup>In units of kcal/mol.

incubation with **11** (5–10 nM final). A 1 nM *humPR3* 262  
 concentration was estimated in these samples by comparison 263  
 with the rate of hydrolysis of the ABZ-VADnVADYQ-EDDnp 264  
 substrate. Cleavage of the *humPR3* substrate was totally 265  
 inhibited after incubation for 20 min at 37 °C with **11** (Figure 266

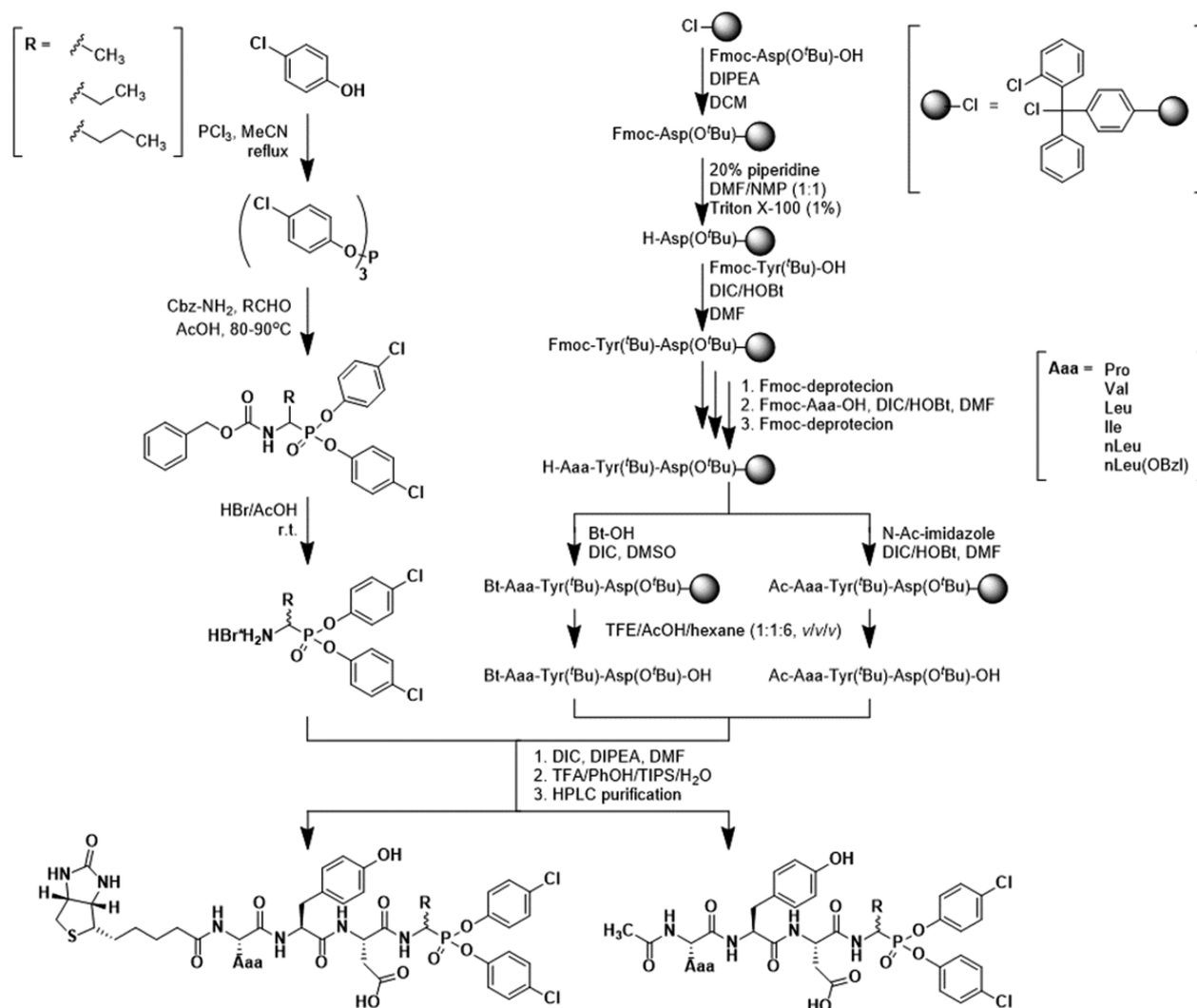


**Figure 4.** Inhibition of *macPR3* by inhibitor **11**. (A) Human and macaque sequence alignment. The sequences of *humPR3* (1FUJ.pdb)<sup>12</sup> and *macPR3* (*Macaca fascicularis*, GenPept: XP\_005587394.1) were aligned using Protein BLAST with default parameters. Similar amino acid residues are indicated in blue and remaining substitutions are in red. Active center residues are indicated by asterisk. The residues included in quantum chemical calculations are indicated in bold. The sequence numbering according 1FUJ.pdb file. Sequence alignment of *humPR3* and *macPR3* show 190/221 (86%) identical positions, 200/221 (90%) positives, and no gaps. (B) Proposed putative model of **11** in *macPR3* active center. (C) Inhibition of PR3 by **11** in macaque neutrophil lysates. The volume of lysates was adjusted to give final concentrations of PR3, 1 nM. Diluted samples were incubated with **11** (10 nM final) for 20 min at 37 °C. Inhibition was monitored by measuring the residual activity of PR3 on ABZ-VADnVADYQ-EDDnp (20 μM) as in Korkmaz et al.<sup>52</sup> Purified *humPR3* (1 nM) was used as control. Similar results were found in three independent experiments.

267 **3B**), while *humNE* activity remained unchanged (not shown).  
 268 Inhibitor **11** was also successfully used to selectively label  
 269 proteolytically active *humPR3* in CF sputum and in a lysate of  
 270 purified human blood neutrophils (Figure 3C). Additionally,  
 271 we showed that inhibitor **11** preserved full inhibitory activity  
 272 and resisted degradation when it was mixed with CF sputum  
 273 for 2 h at 37 °C as shown by high performance liquid  
 274 chromatography (HPLC) (Figure 3D).

275 **Characterization and Inhibition of Macaque PR3**  
 276 (*macPR3*) by Phosphonate Inhibitors in Purified Neu-  
 277 trophil Lysates. Protein sequences alignment of *humPR3* and  
 278 macaque PR3 (*Macaca fascicularis*) shows that they are 86%  
 279 identical, and they differ by only 28 residues. Their substrate  
 280 binding site is very similar and critical residues Lys99, Arg143,

and Ile217 that confer high selectivity to *humPR3* are conserved  
 in *macPR3* (Figure 4A,B). We thus hypothesized that, unlike  
 PR3 homologues in rodents, *macPR3* will be efficiently  
 inhibited by peptide-based phosphonate inhibitors designed  
 for *humPR3*. All phosphonate inhibitors of *humPR3* reported  
 above were able to inhibit *macPR3*. As observed for *humPR3*,  
 biotinylated inhibitors were more efficient than acylated  
 inhibitors at inhibiting *macPR3* (**1**, Ac-PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-  
 Cl)<sub>2</sub>,  $k_{\text{obs}}/I = 55 \pm 4 \text{ M}^{-1} \text{ s}^{-1}$ ; **2**, Bt-PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>,  
 $k_{\text{obs}}/I = 1985 \pm 215 \text{ M}^{-1} \text{ s}^{-1}$ ; **11**, Bt-VYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>,  
 $k_{\text{obs}}/I = 36480 \pm 3350 \text{ M}^{-1} \text{ s}^{-1}$ ), but their overall potency was  
 somewhat lesser than that recorded for *humPR3* (Table 1). The  
 inhibition of *macPR3* with **11** is shown in Figure 3C.



**Figure 5.** Scheme showing the synthesis of peptidyl di(chlorophenyl)-phosphonate ester inhibitors.

294 To further examine binding preferences of **11** against human  
 295 and *macPR3* proteases and interaction energy values between  
 296 Bt-Val4 fragment of the inhibitor and PR3, binding sites were  
 297 compared for particular residues representing S5 binding  
 298 pocket (Table 4). Lys99, the most important residue promoting  
 299 inhibitor binding of *humPR3*, seems to exert also the largest  
 300 influence in terms of the analogous interaction with *macPR3*.  
 301 However, the corresponding binding energy value is less  
 302 significant in the case of *macPR3*–inhibitor complex compared  
 303 to interaction with *humPR3* (–10.0 versus –16.6 kcal/mol;  
 304 Table 4). Another substantial difference in binding energy  
 305 values concerns repulsive interaction due to the presence of  
 306 Phe215 residue. Unfavorable interaction characterizing  
 307 *humPR3*–inhibitor complex (5.5 kcal/mol) amounts to 24.2  
 308 kcal/mol in the corresponding *macPR3* Phe215–inhibitor  
 309 complex (Table 4). The remaining repulsive interactions  
 310 associated with Ile217 and Trp218 are retained in the case of  
 311 *macPR3* inhibition despite the substitution of Trp218 by an  
 312 arginine residue. Interestingly, three out of five substitutions  
 313 that involve PR3 residues in the vicinity of the Bt-Val4 inhibitor  
 314 fragment do not seem to modulate binding potency of **11**  
 315 against human and *macPR3* homologues. The more substantial  
 316 changes related to residue substitution accompany the change  
 317 of Phe166 to leucine and Gly219 to glutamate. However, these

substitution-induced changes in binding energy cancel each  
 318 other out, as the interaction energy value increased by 3 kcal/  
 319 mol as a result of the Phe166Leu substitution is decreased by  
 320 the same extent upon the Gly219Glu substitution. Overall, the  
 321 differences in inhibitor binding by human and *macPR3* appear  
 322 to arise from decreased attractive interaction with Lys99 and  
 323 increased repulsion with Phe215 residues. Because conforma-  
 324 tion and spatial placement of these two residues is essentially  
 325 identical in both complexes, the observed changes in binding  
 326 energy appear to arise from slightly different positioning of the  
 327 Bt-Val4 portion of the inhibitor molecule due to substitutions  
 328 present in the *macPR3* S4 and S5 subsites. 329

## DISCUSSION

Evidence has now accumulated that the neutrophilic serine  
 331 protease *humPR3* acquired specific pathophysiological proper-  
 332 ties and nonredundant functions in spite of its close  
 333 resemblance to *humNE*.<sup>5,29</sup> Indeed, it slightly differs from the  
 334 latter by its spatiotemporal localization,<sup>30</sup> its substrate  
 335 specificity, and its sensitivity to natural inhibitors, all factors  
 336 that taken together explain its specific function as an  
 337 autoantigen in granulomatosis with polyangiitis and its probable  
 338 involvement in cell apoptosis.<sup>6,11</sup> Controlling the proteolytic  
 339 activity of this protease specifically, e.g., by protease inhibitors,  
 340

341 is a means to better understand its biological function, but all  
342 physiological inhibitors of *humPR3* preferentially target  
343 *humNE*. It is only recently that we and others began to  
344 synthesize chemical inhibitors that selectively target the  
345 *humPR3* active site.<sup>7</sup> The specificity of serine proteases is  
346 determined by their substrate binding sites that are located on  
347 both sides of the cleaved peptide bond. We used a substrate-  
348 based approach to develop serpin-like irreversible inhibitor  
349 (SerpinB1(STDA/R) and azapeptide (azapro-3), a reversible  
350 inhibitor that selectively inhibits PR3.<sup>31</sup> Such inhibitors,  
351 however, cannot be used as ABP to visualize active *humPR3*  
352 in biological fluids or in cells and tissues. We recently  
353 developed a series of N-terminally biotinylated peptidyl-  
354 diphenyl phosphonate inhibitors that allow the detection of  
355 *humPR3* at the cell surface and inside cells.<sup>18</sup> These are  
356 transition state analogues, irreversible inhibitors that interact  
357 with nonprime subsites of the target serine protease to form  
358 “phosphorylated” enzymes. Protease–inhibitor complexes  
359 show a remarkable stability due to the similarity of the  
360 phosphorus atom with the tetrahedral intermediate formed  
361 during peptide bond hydrolysis. Although chemically stable in  
362 blood samples, their pharmacological use requires that they  
363 interact rapidly with their target protease to be effective at low  
364 concentrations. We have further investigated the nonprime  
365 specificity of *humPR3* to develop more potent di-  
366 (chlorophenyl)-phosphonate ester inhibitors that could be  
367 used as molecular probes to control *humPR3* activity (Figure  
368 5).

369 We previously showed that the S2/P2 specificity was  
370 essential to discriminate between *humPR3* and its close  
371 homologue *humNE*.<sup>15</sup> Lys99 in *humPR3* is a key residue to  
372 explain the preferential accommodation of negatively charged  
373 or polar residues at P2.<sup>5,7</sup> Thus, selective *humPR3* substrates or  
374 peptide sequences selectively cleaved by *humPR3* all contain a  
375 negatively charged or a polar residue at position P2.<sup>6</sup>  
376 SerpinB1(STDA/R) and azapro-3 that selectively inhibit  
377 *humPR3* contain a negatively charged residue (Asp) at P2  
378 position. However, *humPR3* may accommodate different  
379 residues at P1 and P4 as confirmed by molecular modeling  
380 studies. The S1 binding pocket of *humPR3* is more accessible  
381 and spacey than that of *humNE* and can accommodate not only  
382 the Ala or Abu side chain but also methionine, valine, and nVal,  
383 which was shown experimentally and by computational  
384 docking. In a recent study using single-residue mutant of  
385 *humPR3* with Arg at position 217 (PR3I217R), we showed that  
386 Ile217 located in the neighborhood of the S4 subsite pocket  
387 significantly affects the substrate specificity of *humPR3*.<sup>18</sup> The  
388 docking models performed in this study using phosphonate  
389 inhibitors indicate also that the solvent accessible surface of the  
390 S4 subsite is limited by Trp218 and Ile217 on one side. The  
391 latter two residues are most likely responsible for the binding  
392 preference toward aliphatic side chains at P4 and Lys99, which  
393 is located on the opposite side of the S4 subsite, determines  
394 cooperation between S2 and S4 via hydrogen bonding.  
395 Introduction of a Val and a nVal at P4 and P1 positions,  
396 respectively, in the biotinylated *humPR3* inhibitor previously  
397 reported, Bt-PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (2), enhanced the  $k_{\text{obs}}/[I]$   
398 value toward *humPR3* by ~20-fold. This was probably because  
399 the substitution of Pro4 by Val4 improved the flexibility of the  
400 inhibitor, favoring the formation of hydrogen bonds between  
401 the  $\epsilon$ -amino group of Lys99 and Bt-Val4 amide carbonyl  
402 oxygen as well as Asp2 carboxyl group. These hydrogen bond

interactions are in agreement with previously described  
cooperation observed between S2 and S4 subsites.<sup>18</sup>

Biotin at the N-terminal of P4 residue in phosphonate  
inhibitors and peptidyl-pNA substrates displays stabilizing  
properties. General orientation, size, and hydrophobic character  
of *humPR3* S5 pocket that accommodates N-terminal biotin is  
similar to that of *humNE* crystallized in complex with a  
phosphonate inhibitor bearing a nLeu(O-Bzl) moiety at P4 and  
called an “exopocket”, an extension of the S4 subsite.<sup>32</sup> The  
docking models from this study show that the terminal cavity of  
the *humPR3* S5 pocket formed by Phe166, Cys168, and Arg177  
accommodates the biotin heterocyclic rings, while the hydro-  
phobic surface of Ile217 interacts with the biotin aliphatic  
chain. The location of the biotin rings into the S5 binding site  
prevents recognition of the biotinylated inhibitors by extravidin  
by WB under nondenaturing conditions (not shown). Only  
the five-carbon aliphatic chain of biotin participates in the  
stabilization of the inhibitor within the *humPR3* active site as  
deduced from the observation that a phosphonate inhibitors  
with a same peptide sequence but bearing only a N-terminal  
five-carbon aliphatic chain enhanced the inhibition rate as well  
as whole biotin.<sup>18</sup> Analysis of the docking models suggests that  
the substantial impact of N-terminal biotin binding with S5  
subsite on the overall inhibitory potency might be connected  
with the limited size of S4 binding site. The main contribution  
to the binding energy at this position is provided by Lys99,  
forming the hydrogen bond with the backbone of the inhibitor  
and stabilizing the biotin moiety in proper orientation. Because  
of the narrow character of S4 subsite and the presence of  
Trp218 and Ile217 on the opposite site of Lys99, there is a  
strong preference for small, hydrophobic residues at P4  
position. Therefore, the introduction of more sizable side  
chain such as nLeu(O-Bzl) may influence proper stabilization of  
the compound at P4 subsite, resulting in decreased inhibitory  
potency.

One of the challenges when designing preclinical studies for  
PR3 is to select a relevant animal model. We previously showed  
that PR3 from rodents differs from *humPR3* both in terms of  
substrate specificity, which preclude the use of substrate-derived  
phosphonate inhibitors and of subcellular distribution because  
there is no constitutive expression of PR3 at the neutrophil  
surface of rodent neutrophils.<sup>27,33</sup> We therefore used a  
nonhuman primate model to investigate the substrate  
specificity of neutrophilic PR3 and its sensitivity to  
phosphonate inhibitors developed against *humPR3*. In view of  
the highly conserved primary amino acid sequence of *macPR3*  
implying a very similar specificity as with *humPR3*, *macPR3*  
cleaved the *humPR3* substrate at the same site<sup>34</sup> and this  
activity was inhibited by all phosphonate inhibitors of *humPR3*  
used in this study. WB analysis of the macaque neutrophil  
lysate using an anti-*humPR3* antibody revealed the presence of  
a single band of 26 kDa in the neutrophil lysate with no  
glycosylated forms.<sup>34</sup> A single band of 75 kDa was revealed  
after the lysate was incubated with human  $\alpha$ 1PI, indicating that  
*macPR3* had formed an irreversible complex with the serpin. In  
keeping with this observation, the proteolytic activity toward  
the *humPR3* substrate in the lysate was inhibited by  $\alpha$ 1PI and  
by the PR3-specific serpinB1(STDA/R) inhibitor<sup>35</sup> (not  
shown). The identification of proteolytically active *macPR3* in  
the neutrophil lysate was further confirmed by electrophoresis  
under nondenaturing/nonreducing conditions using the ABP  
Bt-[PEG]66-PYDAP(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> and streptavidin-peroxi-  
dase staining.<sup>34</sup> We found similar level of active PR3 in lysates

466 of purified human and macaque neutrophils by kinetics and  
467 immunoblotting assays. The macaque model thus appears as  
468 relevant animal model for in vivo studies.

## 469 ■ CONCLUSION

470 Targeting the *humPR3* active site by specific inhibitors has  
471 become evidence as soon as it has been established that it was  
472 not a redundant protease mimicking *humNE* and that its  
473 proteolytic activity was poorly controlled by physiological  
474 inhibitors. We have optimized here the structure of peptidyl  
475 phosphonate inhibitors by coupling molecular modeling studies  
476 with kinetic analyses, and we obtained molecular probes to  
477 follow the fate and further investigate the function of PR3 both  
478 in vitro and in vivo. The potency and selectivity of the  
479 inhibitors developed here let us suppose that they are suitable  
480 therapeutic tools for fighting inflammatory and/or infectious  
481 diseases where the role of *humPR3* has been clearly identified  
482 or even only suspected.

## 483 ■ EXPERIMENTAL SECTION

484 **Materials.** *humNE* (EC 3.4.21.37) was obtained from Athens  
485 Research and Technology (USA). The fluorescence resonance energy  
486 transfer (FRET) substrates ABZ-VADnVADYQ-EDDnp/ABZ-  
487 APEEIMRRQ-EDDnp and chromogenic *para*-nitroanilide substrates  
488 synthesized by Genecust (Dudelange Luxembourg). IGEPAL CA-630  
489 (NP40) was purchased from Sigma-Aldrich (St. Louis, MO, USA).

490 **Synthesis of Peptidyl-phosphonate Inhibitors.** All reagents  
491 and solvents were obtained from commercial sources and were used  
492 without purification.

493 All final compounds were purified to >95% purity HPLC system  
494 (Jasco LC System, Jasco, Japan) equipped with Supelco Wide Pore C8  
495 column (8 mm × 250 mm) and ultraviolet-visible (UV-vis, 226 nm)  
496 and fluorescent detectors (excitation 320 nm, emission 450 nm). A  
497 linear gradient from 10 to 90% of B within 40 min was applied (A,  
498 0.1% TFA in water; B, 80% acetonitrile in A).

499 The nuclear magnetic resonance spectra ( $^1\text{H}$ ,  $^{31}\text{P}$  and  $^{13}\text{C}$ ) were  
500 recorded on either a Bruker Avance DRX-300 (300.13 MHz for  $^1\text{H}$   
501 NMR, 121.50 MHz for  $^{31}\text{P}$  NMR), a Bruker Avance 600 MHz (600.58  
502 MHz for  $^1\text{H}$  NMR, 243.10 MHz for  $^{31}\text{P}$  NMR, and 101.12 MHz for  
503  $^{13}\text{C}$  NMR) or Bruker AVANCE III 700 MHz (700.67 MHz for  $^1\text{H}$   
504 NMR) spectrometer. Chemical shifts are reported in parts per million  
505 (ppm) relative to a tetramethylsilane internal standard. Mass spectra  
506 were recorded using a Biflex III MALDI TOF mass spectrometer  
507 (Bruker, Germany). The cyano-4-hydroxycinnamic acid (CCA) was  
508 used as a matrix. High resolution mass spectra (HRMS) were acquired  
509 either on a Waters Acquity Ultra Performance LC, LCT Premier XE,  
510 or Bruker micrOTOF-Q II mass spectrometer.

511 **Cbz-Protected 1-Aminoalkylphosphonate Diaryl Esters (General**  
512 **Procedure).** The first step in the synthesis of the phosphonic  
513 analogues of Ala, nVal, and Abu was the preparation of tri(4-  
514 chlorophenyl)phosphite from 4-chlorophenol and phosphorus tri-  
515 chloride.<sup>36</sup> Briefly, phosphorus trichloride (10 mmol) was added to 4-  
516 chlorophenol (30 mmol) dissolved in acetonitrile (50 mL) and the  
517 mixture refluxed for 6 h. The volatile elements were removed in a  
518 vacuum, and the resulting crude phosphite, a yellow oil, was used  
519 directly in an amidalkylation reaction. It was mixed with benzyl  
520 carbamate (12 mmol) and an appropriate aldehyde: acetaldehyde,  
521 butyraldehyde, or propionaldehyde (12 mmol) and refluxed in acetic  
522 acid for 3 h (Oleksyszyn's method<sup>37</sup>).

523 **Deprotection of Cbz Group (General Procedure).** The Cbz  
524 protecting group was removed by incubation with 33% hydrobromic  
525 acid in acetic acid (2 h). The volatile components were removed under  
526 reduced pressure, and the products were crystallized from methanol/  
527 diethyl ether to give target compounds as hydrobromide salts.

528 **Benzyl (1-(Bis(4-chlorophenoxy)phosphoryl)ethyl)carbamate (12,**  
529 **Cbz-Ala<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>).** 12 was prepared using the general method  
530 described above and crystallized from methanol to yield a white solid

(56%).  $^1\text{H}$  NMR (300.13 MHz,  $\text{CDCl}_3$ -*d*<sub>1</sub>, ppm):  $\delta$  7.43–6.97 (m, 531  
14H), 5.22–5.08 (m, 2H), 4.74–4.37 (m, 1H), 1.56 (dd, *J* = 18.2, 7.4 532  
Hz, 3H).  $^{31}\text{P}$  NMR (121.50 MHz,  $\text{CDCl}_3$ -*d*<sub>1</sub>, ppm):  $\delta$  19.56 (s).  $^{13}\text{C}$  533  
NMR (101.12 MHz,  $\text{DMSO-}d_6$ , ppm):  $\delta$  156.16, 156.11, 151.14, 534  
151.05, 150.84, 150.74, 137.20, 134.14, 131.91, 131.88, 128.87, 128.46, 535  
128.34, 126.15, 126.03, 121.13, 121.09, 120.99, 119.99, 119.73, 66.48, 536  
45.30, 43.73, 15.52. HRMS: calcd for  $(\text{C}_{22}\text{H}_{20}\text{Cl}_2\text{NO}_3\text{P})\text{H}^+$  480.0534, 537  
found 480.0533. 538

**Benzyl (1-(Bis(4-chlorophenoxy)phosphoryl)propyl)carbamate** 539  
**(13, Cbz-Abu<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>).** 13 was prepared using the general 540  
method described above and crystallized from methanol to yield a 541  
white solid (19%).  $^1\text{H}$  NMR (300.13 MHz,  $\text{CDCl}_3$ -*d*<sub>1</sub>, ppm):  $\delta$  7.45– 542  
6.99 (m, 13H), 5.14 (d, *J* = 10.7 Hz, 1H), 5.24–5.06 (m, 2H), 4.50– 543  
4.33 (m, 1H), 2.20–2.03 (m, 1H), 1.87–1.64 (m, 1H), 1.11 (t, *J* = 7.3 544  
Hz, 3H).  $^{31}\text{P}$  NMR (121.50 MHz,  $\text{CDCl}_3$ -*d*<sub>1</sub>, ppm):  $\delta$  18.31 (s).  $^{13}\text{C}$  545  
NMR (101.12 MHz,  $\text{DMSO-}d_6$ , ppm):  $\delta$  156.85, 156.81, 149.45, 546  
149.35, 149.12, 149.03, 137.34, 130.35, 130.28, 130.03, 129.86, 129.65, 547  
128.88, 128.44, 128.32, 128.01, 122.94, 122.90, 122.67, 122.63, 117.44, 548  
66.43, 51.27, 49.72, 22.46, 22.42, 11.16, 11.01. HRMS: calcd for 549  
 $(\text{C}_{23}\text{H}_{22}\text{Cl}_2\text{NO}_3\text{P})\text{H}^+$  494.0691, found 494.0699. 550

**Benzyl (1-(Bis(4-chlorophenoxy)phosphoryl)butyl)carbamate** 551  
**(14, Cbz-nVal<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>).** 14 was prepared using the general 552  
method described above and crystallized from methanol to yield a 553  
white solid (20%).  $^1\text{H}$  NMR (600.58 MHz,  $\text{CDCl}_3$ -*d*<sub>1</sub>, ppm):  $\delta$  7.42– 554  
6.68 (m, 13H), 5.24–5.15 (m, 2H), 5.11 (d, *J* = 12.2 Hz, 1H), 4.57– 555  
4.44 (m, 1H), 2.07–1.95 (m, 2H), 1.67–1.39 (m, 2H), 1.02–0.93 (m, 556  
3H).  $^{31}\text{P}$  NMR (243 MHz,  $\text{CDCl}_3$ , ppm):  $\delta$  18.47 (s).  $^{13}\text{C}$  NMR 557  
(101.12 MHz,  $\text{DMSO-}d_6$ , ppm):  $\delta$  156.85, 156.76, 156.71, 149.45, 558  
149.35, 149.11, 149.02, 137.32, 130.36, 130.29, 130.05, 129.87, 129.65, 559  
128.88, 128.45, 128.34, 122.95, 122.91, 122.67, 122.63, 117.44, 66.44, 560  
49.23, 47.66, 30.63, 19.10, 18.95, 13.68. HRMS: calcd for 561  
 $(\text{C}_{24}\text{H}_{24}\text{Cl}_2\text{NO}_3\text{P})\text{Na}^+$  530.0667, found 530.0670. 562

**Bis(4-chlorophenyl) (1-Aminoethyl)phosphonate Hydrobromide** 563  
**(15, HBr×H<sub>2</sub>N-Ala<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>).** 15 was prepared using the general 564  
method described above and crystallized from diethyl ether to yield a 565  
white solid (97%).  $^1\text{H}$  NMR (300.13 MHz,  $\text{DMSO-}d_6$ , ppm):  $\delta$  8.85 566  
(s, 3H), 7.57–7.44 (m, 4H), 7.36–7.16 (m, 4H), 4.45–4.24 (m, 1H), 567  
1.55 (dd, *J* = 18.3, 7.2 Hz, 3H).  $^{31}\text{P}$  NMR (121.50 MHz,  $\text{DMSO-}d_6$  568  
ppm):  $\delta$  16.49 (s).  $^{13}\text{C}$  NMR (101.12 MHz,  $\text{DMSO-}d_6$ , ppm):  $\delta$  569  
148.63, 148.61, 148.53, 148.52, 130.57, 130.54, 130.53, 129.62, 123.02, 570  
122.98, 122.94, 117.45, 43.50, 41.93, 13.96, 13.93. HRMS: calcd for 571  
 $(\text{C}_{14}\text{H}_{14}\text{Cl}_2\text{NO}_3\text{P})\text{H}^+$  346.0167, found 346.0172. 572

**Bis(4-chlorophenyl) (1-Aminopropyl)phosphonate Hydrobro-** 573  
**midate (16, HBr×H<sub>2</sub>N-Abu<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>).** 16 was prepared using 574  
the general method described above and crystallized form diethyl ether 575  
to yield a white solid (83%).  $^1\text{H}$  NMR (300.13 MHz,  $\text{DMSO-}d_6$  576  
ppm):  $\delta$  8.87 (s, 2H), 7.67–7.13 (m, 8H), 4.21 (dt, *J* = 13.6, 6.8 Hz, 577  
1H), 2.24–1.74 (m, 2H), 1.11 (t, *J* = 7.4 Hz, 3H).  $^{31}\text{P}$  NMR (121.50 578  
MHz,  $\text{DMSO-}d_6$ , ppm):  $\delta$  16.50 (s).  $^{13}\text{C}$  NMR (101.12 MHz,  $\text{DMSO-}$  579  
 $d_6$ , ppm):  $\delta$  148.56, 148.47, 130.58, 130.54, 123.05, 123.01, 122.97, 580  
122.93, 48.80, 47.26, 22.01, 21.99, 10.81, 10.72. HRMS: calcd for 581  
 $(\text{C}_{15}\text{H}_{16}\text{Cl}_2\text{NO}_3\text{P})\text{H}^+$  360.0318, found 361.1123. 582

**Bis(4-chlorophenyl) (1-Aminobutyl)phosphonate hydrobromide** 583  
**(17, HBr×H<sub>2</sub>N-nVal<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>).** 17 was prepared using the 584  
general method described above and crystallized from diethyl ether 585  
to yield a white solid (89%).  $^1\text{H}$  NMR (300.13 MHz,  $\text{DMSO-}d_6$  586  
ppm):  $\delta$  8.88 (s, 2H), 7.49–7.27 (m, 8H), 4.25 (dt, *J* = 13.9, 7.0 Hz, 587  
1H), 2.06–1.78 (m, 2H), 1.71–1.46 (m, 2H), 0.90 (t, *J* = 7.3 Hz, 3H). 588  
 $^{31}\text{P}$  NMR (121.50 MHz,  $\text{DMSO-}d_6$ , ppm):  $\delta$  16.55 (s).  $^{13}\text{C}$  NMR 589  
(101.12 MHz,  $\text{DMSO-}d_6$ , ppm):  $\delta$  148.59, 148.58, 148.49, 148.48, 590  
130.58, 130.53, 123.06, 123.02, 122.96, 122.92, 47.40, 45.85, 30.39, 591  
30.36, 19.02, 18.92, 14.06. HRMS: calcd for  $(\text{C}_{16}\text{H}_{18}\text{Cl}_2\text{NO}_3\text{P})\text{H}^+$  592  
374.0474, found 375.1931. 593

The peptides were synthesized manually by the solid-phase method 594  
using Fmoc chemistry. The following amino acid derivatives were 595  
used: Fmoc-Pro, Fmoc-Val, Fmoc-Leu, Fmoc-Ile, Fmoc-nLeu, Fmoc- 596  
nLeu(O-Bzl), Fmoc-Tyr(tBu), and Fmoc-Asp(OtBu). The protected 597  
derivative of the C-terminal amino acid residue, Fmoc-Asp(OtBu), was 598  
attached to the 2-chlorotrityl resin (substitution of Cl 1.46 mequiv/g) 599  
(Calbiochem-Novabiochem AG, Switzerland) in the presence of an 600

601 equimolar amount of diisopropylethylamine (DIPEA) under anhy-  
 602 drous conditions in dichloromethane (DCM) solution. A peptide  
 603 chain was elongated in consecutive cycles of deprotection (20%  
 604 piperidine in dimethylformamide (DMF)/*n*-methylpyrrolidone  
 605 (NMP) (1:1, v/v) with 1% Triton X-100) and coupling (DIC/  
 606 HOBT chemistry; 3 equiv of protected amino acid derivatives were  
 607 used). A 10-fold molar excess of *N*-acetylimidazole in DMF was used  
 608 for acetylation of the N-terminus. Bt-[PEG]<sub>2</sub>-Pro-Tyr-Asp-Ala<sup>P</sup>(O-  
 609 C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> was synthesized via coupling of the Fmoc-PEG<sub>2</sub> to the  
 610 amino group of terminal Pro residue. The N-terminal biotin group was  
 611 conjugated using a 5-fold molar excess of biotin and 1,3-  
 612 diisopropylcarbodiimide (DIC) as the coupling agent in anhydrous  
 613 DMSO for 6 h at 30 °C. The synthesized peptides were cleaved from  
 614 the resin with TFE/hexane/acetic acid (1.6:1, v/v/v).

615 Fully protected peptides were dissolved in DMF and their carboxyl  
 616 groups were activated with DIC and coupled with HBr×H<sub>2</sub>N-Ala<sup>P</sup>(O-  
 617 C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>, HBr×H<sub>2</sub>N-Abu<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>, or HBr×H<sub>2</sub>N-  
 618 nVal<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> in DMF in the presence of DIPEA. The mixture  
 619 was stirred for 6 h, and the DMF was removed under reduced  
 620 pressure. The resulting compounds were suspended in trifluoroacetic  
 621 acid (TFA)/phenol/triisopropylsilane/H<sub>2</sub>O (88:5:2:5, v/v/v/v) for 2  
 622 h to remove side chain protecting groups.

623 The crude peptides were purified by HPLC on a Beckman Gold  
 624 System (Beckman, USA) with an RP Kromasil-100, C8, 5 μm column  
 625 (8 mm × 250 mm) (Knauer, Germany). The solvent systems were  
 626 0.1% TFA (A) and 80% acetonitrile in A (B). Either isocratic  
 627 conditions or a linear gradient were applied (flow rate 3.0 mL/min,  
 628 monitored at 226 nm). The purity of the synthesized peptides was  
 629 verified on RP Kromasil 100, C8, 5 μm column (4.6 mm × 250 mm)  
 630 (Knauer, Germany). The peptides were eluted with a linear gradient of  
 631 the above solvent system (10%–90% B) for 30 min, flow rate 1 mL/  
 632 min, monitored at 226 nm. HPLC retention times and <sup>1</sup>H NMR  
 633 spectra of final phosphonate peptide inhibitors are shown in Table 5

the method of Tian and Tsou.<sup>39</sup> Product formation in the presence of  
 an irreversible inhibitor approaches an asymptote in this system, as  
 described by  $\log([P_{\infty}] - [P]) = \log[P_{\infty}] - 0.43A[Y]t$ .

- where  $[P_{\infty}]$  is the concentration of product formed at time approaching infinity,  $[P]$  is the concentration of product at time  $t$ ,  $[Y]$  is the inhibitor concentration, and  $A$  is the apparent inhibition rate constant in the presence of the substrate.  $A$  is given by  $A = k_{+o}/(1 + K^{-1}[S])$
- where  $k_{+o}$  is the rate constant for association of the inhibitor with the enzyme,  $K^{-1}$  is the inverted Michaelis constant, and  $[S]$  the substrate concentration. The apparent inhibition rate constant  $A$  is the slope of a plot of  $\log([P_{\infty}] - [P])$  against  $t$ , to give the second-order rate constant of inhibition  $k_{+o}$ .

The rates of inhibition of purified *humPR3*, *macPR3* (in purified blood neutrophil lysates), and purified *humNE* were measured using FRET substrates (ABZ-VADnVADYQ-EDDnp (10 μM final) and ABZ-APEEMRRQ-EDDnp (10 μM final) in 50 mM HEPES, 0.75 M NaCl, and 0.05% NP40, pH = 7.4; excitation wavelength, 320 nm; emission wavelength, 420 nm; Spectramax Gemini (Molecular Devices, Sunnyvale, CA, USA). Final protease concentrations were 1 nM.

$K_i$  and  $k_2$  determination: We monitored the extent of protease inhibition at several time points for a different inhibitor concentrations  $[I]$ . The observed rate constant for inhibition,  $k_{obs}$ , at each concentration was determined from the slope of a semilogarithmic plot of inhibition versus time. The  $k_{obs}$  values were replotted against inhibitor concentration and fitted to a hyperbolic equation,  $k_{obs} = k_2[I]/(K_i + [I])$ , to obtain values for  $K_i$  and  $k_2$ .<sup>40</sup>

**Detection of PR3 in Biological Fluids.** CF sputa (50 μg proteins) were incubated with **11** (50 nM final) for 20 min at 37 °C in PBS. The reaction was stopped by adding 1 volume of 2× SDS reducing buffer and heating at 90 °C for 5 min. The components of the mixture were separated by SDS-PAGE, 12% NaDodSO<sub>4</sub>-polyacrylamide gel electrophoresis under denaturing conditions. They were transferred to a nitrocellulose (Hybond)-ECL (Enhanced Chemiluminescence) membrane at 4 °C.

**Extravidin Peroxidase Detection.** Free sites on the membrane were blocked with 3% bovine serum albumin (BSA) in 0.1% Tween in PBS for 90 min at room temperature (RT). Membranes were then given two quick washes with PBS-Tween 0.1% and incubated for 2 h at RT with extravidin horseradish peroxidase (HRP) (Sigma-Aldrich) (diluted 1/4000 in 3% BSA in PBS-Tween 0.1%). The extravidin-HRP treated membrane was washed (3 × 10 min) with PBS-Tween 1% and then incubated with HRP substrate for 3 min. Reactive bands were identified by chemiluminescence (ECL Kit).

**Immunodetection.** Free sites on the membranes were blocked by incubation with 5% nonfat dried milk in PBS-0.1% Tween for 90 min at RT. They were washed twice with PBS-Tween 0.1% and incubated overnight with a rabbit primary anti-PR3 antibody (1:700, EPR6277 Abcam), followed by a goat antirabbit IgG secondary antibody (1:7000, A9169 Sigma). These membranes were then washed and processed as above.

**Purification and Lysis of *M. fascicularis* Neutrophils.** Female cynomolgus monkeys (*Macaca fascicularis*) (approximately 3 years old and weighing 4–5 kg) were obtained from a commercial supplier. All animal experiments and procedures were approved by the local animal experimentation ethics committee (Comité d'éthique Val de Loire (APAFIS no. 2982-20151105293399v6)). Five mL of peripheral blood samples were collected in lithium–heparin tubes from a femoral vein. Animals were kept under spontaneous ventilation during anesthesia with ketamine (10 mg/kg). The monitoring included pulse-oximetry and heart rate recording. Intravenous access was secured with a 22G canula on the legs. Anticoagulated whole blood was layered onto Ficoll density gradient and centrifuged. The purified neutrophils (>98%) in suspension was treated with H<sub>2</sub>O for 30 s to lyse red blood cells. The neutrophils were then lysed in Hepes 50 mM, NaCl 0.15 M, NP40 0.5%, pH 7.4, and the supernatant was collected and stored at –80 °C.

**Chromatographic Procedures and Peptide Analysis.** Inhibitor **11** (75 μM final) was incubated with the cell free supernatants of sputa

**Table 5. Calculated and Observed Masses<sup>a</sup> and HPLC Retention Times of Synthesized Inhibitors 1–11**

compd	calculated mass (Da)	found mass (Da)	retention time (min)
1	763.56	764.67	12.36
2	947.82	948.91	13.12
3	1266.18	1266.23	10.05
4	949.83	951.01	12.56
5	963.86	964.79	12.47
6	963.86	964.92	12.51
7	963.86	964.88	12.42
8	1055.95	1057.08	13.57
9	961.54	962.50	13.43
10	975.87	976.95	13.20
11	977.89	977.97	12.58

<sup>a</sup>The obtained molecular weights represent pseudomolecular ions (M + H)<sup>+</sup>.

and Supporting Information, respectively. Mass spectrometric analysis of the inhibitors (Table 5) was done on a MALDI MS (a Biflex III MALDI-TOF spectrometer, Bruker Daltonics, Germany) using a CCA matrix.

**Enzymatic Studies: Free *humPR3* and *humNE* Were Titrated with α1PI.**  $k_{cat}/K_m$  determination: The specificity constants  $k_{cat}/K_m$  for peptidyl-pNA substrates were determined under first-order conditions.<sup>38</sup> The cleavage of the substrates (1 mM final) was monitored by measuring the absorbance of liberated pNA at 410 nm on the spectrophotometer (Versamax microplate reader, Molecular Devices, Sunnyvale, CA, USA). Measurements were carried out at 37 °C in buffer 50 mM HEPES, 0.75 M NaCl, 0.05% NP40, pH 7.4. Final protease concentrations were 0.01–1 μM.

$k_{obs}/[I]$  determination: The inactivation of proteases by phosphonate inhibitors (substrate analogue inhibitors) in the presence of the substrate by competition for the enzyme-binding site was measured by

719 from CF patients at 37 °C for 2 h in PBS. FRET substrate ABZ-  
720 VADnVADYQ-EDDnp<sup>15</sup> (20 μM final) was incubated with *humR3*  
721 and macaque neutrophil lysate supernatant (10–500 nM) at 37 °C in  
722 50 mM HEPES, 0.75 M NaCl, and 0.05% NP40, pH = 7.4. The  
723 proteins were precipitated with absolute ethanol (4 volumes). The  
724 supernatant containing the peptides were dried under vacuum and  
725 dissolved in 200 μL of 0.01% trifluoroacetic acid (v/v), then  
726 fractionated by Agilent Technology 1200 series HPLC system (Agilent  
727 Technology, CA, USA) on a C18 column (2.1 mm × 30 mm, Merck  
728 Millipore) at a flow rate of 0.3 mL/min with a linear gradient (0–90%,  
729 v/v) of acetonitrile in 0.01% trifluoroacetic acid over 40 min. Eluted  
730 peaks were monitored at 220 nm.

731 **Molecular Modeling.** Molecular docking was performed in order  
732 to explain interactions of Ac-PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (**1**), Bt-  
733 PYDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (**2**), Bt-nLeu(O-Bzl)YDA<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub>  
734 (**8**), and Bt-VYDnV<sup>P</sup>(O-C<sub>6</sub>H<sub>4</sub>-4-Cl)<sub>2</sub> (**11**) with *humPR3* and  
735 *macPR3*. As a receptor, the crystal structure of *humPR3* (1FUJ.pdb)<sup>12</sup>  
736 was selected. The same structure was used as a template for *macPR3*  
737 3D model obtained by means of automated homology modeling  
738 server, SWISS-MODEL.<sup>41</sup> For the docking studies, inhibitor molecules  
739 were used as a peptidyl phosphonic acids [Ac-PYDA<sup>P</sup>(OH)<sub>2</sub>, Bt-  
740 PYDA<sup>P</sup>(OH)<sub>2</sub>, Bt-VYDnV<sup>P</sup>(OH)<sub>2</sub>, and Bt-nLeu(O-Bzl)YDA<sup>P</sup>(OH)<sub>2</sub>]  
741 instead of di(chlorophenyl)-phosphonate esters, as this is the form  
742 present in the “aged” protein–inhibitor complex. The ligand models  
743 were optimized using the MM2 force field (as implemented in  
744 ChemBio3D 12.0),<sup>42</sup> while the atom types and protonation of all  
745 structures were set using SPORES.<sup>43</sup> The docking was carried out  
746 using the Protein–Ligand ANT System (PLANTS v. 1.2) with  
747 PLANTS<sub>CHEMPLP</sub> scoring function.<sup>44–46</sup> The protein molecules were  
748 treated as fixed with the binding site center defined at a carbonyl  
749 oxygen of Ser214 and the binding site radius of 15 Å. The distance  
750 constraints were set up to increase the preference of interaction  
751 between (a) inhibitor phosphorus atom and the hydroxide oxygen of  
752 protease Ser195 (distance range was defined between 2.2 and 4.0 Å),  
753 (b) the terminal carbon of Ala/nVal side chain of ligand P1 position  
754 and enzyme S1 binding pocket set at γ-carbon of Ile190 (distance  
755 range: 5.5–6.5 Å for Ala and 2.2–5.0 Å for nVal), (c) Asp γ-carbon of  
756 the inhibitor (P2 position) and PR3 ε-amine nitrogen of Lys99  
757 (distance range 2.0–5.0 Å), and (d) ligand P3–P4 amide bond  
758 nitrogen and Val216 carbonyl oxygen of the receptor (distance range  
759 2.0–5.0 Å). The lowest energy binding poses obtained from docking  
760 simulations were then employed in quantum chemical calculations of  
761 interaction energy between PR3 amino acid residues and Bt-Val4  
762 fragment of inhibitor to explain the differences in activity of **11** toward  
763 human and macaque enzyme. *humPR3* or *macPR3* binding site was  
764 represented by all amino acid residues within 6 Å of inhibitor fragment  
765 considered herein. Because of the presence of disulfide bridge in the  
766 vicinity of inhibitor molecule, covalently linked Cys168 and Cys182  
767 residues were included as a single monomer. Arg177 was found to be  
768 hydrogen-bonded to Asn98 and Asn180 residues. To avoid disrupting  
769 the hydrogen bonding network, these three residues were also  
770 considered as a monomer. The remaining 15 PR3 residues were  
771 included separately. The dangling bonds resulting from cutting the  
772 residues out of the protein scaffold were saturated with hydrogen  
773 atoms. PR3–inhibitor binding energy was calculated in a pairwise  
774 manner at the second-order Møller–Plesset level of theory (MP2)  
775 using 6-31+G(d) basis set<sup>47–49</sup> and counterpoise correction to  
776 eliminate basis set superposition error.<sup>50</sup> Quantum chemical  
777 calculations were performed in Gaussian09 program.<sup>51</sup>

## 778 ■ ASSOCIATED CONTENT

### 779 ● Supporting Information

780 The Supporting Information is available free of charge on the  
781 ACS Publications website at DOI: 10.1021/acs.jmed-  
782 chem.7b01416.

783 Spectroscopic data of synthesized inhibitors; supporting  
784 Information includes <sup>1</sup>H NMR spectra together with  
785 SMILES for compounds **1–11** (PDF)

Docking poses for **1**, **2**, **8**, and **11** with *humPR3* and **11** 786  
with *macR3* (ZIP) 787  
Compound data (CSV) 788

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807 wrote the manuscript. All authors contributed to the writing  
808 and revision processes of the manuscript.

### 809 Notes

The authors declare no competing financial interest. 810

## 811 ■ ACKNOWLEDGMENTS

812 This work was supported by the Ministère de l'Enseignement  
813 Supérieur et de la Recherche, the Région Centre and the Fonds  
814 Européen de Développement Régional (Project INFINHI),  
815 and the Polish Ministry of Science and Higher Education  
816 granted to M.W. (project no. IP2012 0596 72). B.K.  
817 acknowledges the Association Vaincre La Mucoviscidose  
818 (AVLM) and the Alexandre von Humboldt Foundation. M.S.  
819 and R.G. are grateful for support to Wroclaw University of  
820 Science and Technology (statute funds 10401/0194/17). We  
821 thank Lise Vanderlynden (INSERM U-1100) and Heike  
822 Reimann (Comprehensive Pneumology Center, Institute of  
823 Lung Biology and Disease, iLBD) for technical assistance.

## 824 ■ ABBREVIATIONS USED

825 ABP, activity-based probe; α1PI, alpha-1-proteinase inhibitor;  
826 ABZ, *ortho*-aminobenzoic acid; Bt, biotin; HPLC, high  
827 performance liquid chromatography; CG, cathepsin G; CF,  
828 cystic fibrosis; EDDnp, *N*-(2,4-dinitrophenyl)ethylenediamine;  
829 FRET, fluorescence resonance energy transfer; GPA, gran-  
830 ulomatosis with polyangiitis; hum, human; NE, neutrophil  
831 elastase; NSP, neutrophil serine protease; PBS, phosphate-  
832 buffered saline; PEG, polyethylene glycol; PMN, polymorpho-  
833 nuclear neutrophil; pNA, *para*-nitroaniline; PR3, proteinase 3;  
834 WB, Western blot

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