

Thiol isomerases orchestrate thrombosis and haemostasis

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22 SUMMARY

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Since protein disulphide isomerase (PDI) was first described in 1963, researchers have shown 24 conclusively that PDI and sibling proteins are quintessential for thrombus formation. PDI, 25 ERp5, ERp57 and ERp72, which in most cells are located in the endoplasmic reticulum and 26 function to assist the folding of nascent protein, are released from platelets and vascular cells 27 and interact with integrin α IIb β 3 on the outer surface of platelets. At the cell surface they 28 continue to influence protein folding and function, propagating thrombosis and maintaining 29 30 haemostasis. TMX1, which is a transmembrane thiol isomerase, is the first family member shown to negatively regulate platelets known to date. Targets of thiol isomerases have been 31 indentified including integrin a2\beta1, Von Willebrand Factor (VWF), GpIba, Nox-1, Nox-2 32 and tissue factor, all of which are pro-thrombotic, and several of which are on the cell 33 surface. In spite of this, PDI can paradoxically catalyse the delivery of nitric oxide to 34 platelets, which inhibits their function and decreases thrombus formation. Although the 35 overall effect of PDI is to positively regulate platelet activation, it is still unclear how thiol 36 37 isomerases function in pro-thrombotic states, such as obesity, diabetes and cancer. In parallel, there has been a surge in the development of novel thiol isomerase inhibitors, which display 38 39 selectivity, potency and modulate thrombosis and haemostasis. The availability of selective thiol isomerase inhibitors has culminated in clinical trials with promising outcomes for the 40 prevention of cancer-associated thrombosis. Altogether, thiol isomerases are perceived as an 41 orchestrating force that regulates thrombus development. In the current review we will 42 explore the history of PDI in cardiovascular biology, detail known mechanisms of action and 43 summarise known thiol isomerase inhibitors. 44

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KEYWORDS: Protein disulphide isomerase, platelet, thrombosis, thiol isomerase, history,
 inhibitors

48 **1. INTRODUCTION**

Platelets were first discovered in the 19th century as 'small plates' derived from either neutrophils or red blood cells ¹. Interests in platelets were rekindled in the 1960s, almost 100 years after their initial description, when several groups began to investigate their functions ²⁻ ⁴. These cells are regarded as anucleated fractions of megakaryocytes and great scientific effort has been employed to understand their vital task: to prevent and stop mammals from bleeding ⁵ – which makes platelets a remarkable evolutionary adaptation necessary for human survival (reviewed in ⁶).

56 In order to maintain haemostasis, platelets rely on several molecules to either inhibit (e.g. nitric oxide, prostacylin) or induce (e.g. thrombin, thromboxane, etc) platelet activation 57 ⁷. Thrombus development initiates upon exposure of sub-endothelial extracellular matrix 58 proteins following blood vessel injury, and particularly collagens to which von Willebrand 59 factor (VWF) binds. Platelet receptors GPVI and integrin a2B1 bind to collagen, while 60 glycoprotein (GP) Ib-V-IX binds to VWF^{8,9}. Together these receptors start intracellular 61 (inside-out) signalling that culminates in conformational changes in integrin aIIbβ3, which 62 initiates ligand binding. Fibrinogen and VWF then bind to the integrin, creating a second 63 wave of signalling events, termed outside-in signalling (reviewed in ¹⁰). This latter chain of 64 65 signals created by integrin aIIb₃ binding and clustering results in irreversible platelet adhesion, aggregation, pseudopodia formation and reinforces degranulation of dense- and a-66 granules. In this way key activatory molecules such as ADP and serotonin are secreted, while 67 metabolism of arachidonic acid liberated from platelet membranes results in the production of 68 thromboxane A2 (TxA₂). Through the actions of these secondary activators on platelet 69 receptors, a positive feedback loop is initiated, resulting in the activation and recruitment of 70 71 further platelets and the growth of a platelet thrombus, or haemostatic plug, to stem the loss of blood (reviewed in ¹¹). A summary of inside-out and outside-in signalling is presented in 72 Figure 1. 73

Platelet activation and endothelial damage have been shown to result in the release of a 74 number of normally endoplasmic reticulum (ER)-resident proteins into the blood ^{12,13}. These 75 include members of the thiol isomerase family of oxidoreductase enzymes, namely protein 76 disulphide isomerase A1 (herein referred to as PDI), ERp5, ERp57, ERp72 and TMX1 77 (negative regulator), which were previously thought to be restricted to the ER ¹²⁻¹⁵. PDI is the 78 prototype of the thiol isomerase family, also known as thioredoxins, which catalyse 79 reduction, oxidation and isomerisation of disulphide bonds, as well as nitric oxide (NO) 80 transfer through transitrosation (Figure 2)¹⁶. PDI is the product of the P4HB gene, with a 81

molecular mass of 57 kDa and comprises five modules: two thioredoxin-like domains (a and 82 a') that catalyse oxidoreductase reactions, two substrate-binding domains (b and b'), a C-83 terminal extension domain and a cross-linker sequence between b' and a'. Upon activation 84 soluble, non trans-membrane thiol isomerases (PDI, ERp5, ERp57 and ERp72) are released 85 from platelets whereupon they bind to the outer surface of the plasma membrane, being 86 important for platelet function both in vitro and in vivo and supporting thrombosis and 87 haemostasis ^{12,17}. Indeed, extracellular disulphide exchange reactions between thiol 88 isomerases mentioned above and integrin α IIb β 3 regulate integrin activation ^{12,17-20}. 89

In this review we will explore the contribution of PDI family proteins to thrombus development in health and disease. We will draw a historical landscape of key discoveries in the field followed by a description of known targets of PDI that regulate thrombosis. Finally, we provide a comprehensive table of thiol isomerase inhibitors and future perspectives.

94

95 2. HISTORICAL LANDMARKS

96 2.1. Discovery of PDI

97 In 1963, two independent groups made pivotal discoveries of an enzyme that catalysed 98 the reactivation of reduced ribonuclease. The first group, led by Brunó Straub, described that pigeon and chicken pancreas contained a heat-labile 'factor' that induced the reoxidation of 99 reduced RNAse²¹, whereas the second, led by Nobel prize-winning Christian B Anfinsen, 100 made similar observations using a microsomal system from rat livers ²². However, it was only 101 10 years later that PDI was officially named ²³. PDI was also described as an 'insulin 102 protease' since it catalysed the reduction of insulin, in an assay that has been widely 103 disseminated in the field and still used in contemporary work ²⁴. The protein sequence and 104 identification of the CGHC active sites of rat PDI were only performed in the 1980s ²⁵. From 105 hereon, other PDIs were discovered and their relevance to various physiological and 106 pathophysiological processes in cardiovascular cells began to be explored. The historical 107 108 landmarks of the involvement of PDI in thrombosis are summarized in Figure 3.

109 2.2. PDI is found on the outer surface of cardiovascular cells

Initially thought to be restricted to the ER due to a KDEL sequence, PDI was found to be secreted from activated platelets over 30 years after initial reports by Straub's and Afinsen's groups ¹⁴. Subsequently, Essex et al ²⁶ demonstrated that PDI is localized to the external surface of the platelet plasma membrane. In addition, this group has shown that the

majority of platelet PDI is localized on the platelet surface, while other blood cells showed 114 little PDI when compared to platelets. Further work conducted by this group demonstrated 115 that PDI is recruited to the surface of platelets upon activation ²⁷ and while this process has 116 been shown to be dependent on the reorganisation of the actin cytoskeleton, given its 117 intracellular localisation to the dense tubular system of platelets ²⁸ the exact mechanism of 118 release is unclear. Indeed, work conducted by our group ²⁹ and Raturi et al ³⁰ have identified 119 PDI in plasma extracellular vesicles (EVs). These EVs were capable of potentiating platelet 120 aggregation and displayed reductase activity, both of which were inhibited in the presence of 121 122 a functional anti-PDI antibody ³⁰.

PDI is also present on the surface of endothelial cells, and the secretion of endothelial 123 cell-derived PDI is able to modulate thrombospondin-1 activity ³¹. Upon endothelial lesion, 124 PDI is secreted from endothelial cells to potentiate thrombus formation in vivo ¹³. PDI has 125 also been detected on the surface of vascular smooth muscle cells (VSMC) ³² and shown to 126 regulate VSMC migration, differentiation and redox homeostasis ^{33,34}. Therefore, in spite of 127 initial reports describing an ER-resident chaperone with oxidoreductase activity, the 128 relevance of extracellular PDI, also termed peri/epicellular PDI, has been of great interest to 129 the homeostasis of cardiovascular cells. Importantly, Cho et al ³⁵ were the first to demonstrate 130 that peri/epicellular PDI is critical to thrombus formation in a rodent model of thrombosis in 131 vivo. Several groups have reiterated this finding ^{13,36,37}. However, PDI inhibition ²⁷ or 132 genetically deletion ¹⁷ does not completely abrogate platelet responses, suggesting that there 133 are other thiol isomerases on the surface of platelets that may also regulate 134 thromboinflammatory responses. 135

136 2.3. Other thiol isomerases regulate thromboinflammation

The observation that PDI inhibition was unable to completely inhibit platelet responses 137 has allowed the speculation that other thiol isomerases could be expressed on the platelet 138 surface. Indeed, several additional thiol isomerases have been identified on the platelet 139 membrane ^{12,38-40}. These include ERp5, which is recruited to the platelet outer membrane 140 where it binds to integrin $\beta 3^{-18}$. Selective inhibition of ERp5 and/or PDI with selective 141 142 antibodies revealed additive inhibitory effects ¹⁸; findings that were corroborated by Passam et al ⁴¹ who showed that ERp5 is required for thrombus formation *in vivo* and directly binds 143 to integrin β 3. Other thiol isomerases that have been reported in and on mouse and human 144 platelets and megakaryocytes include: ERp57^{29,38}, ERp72^{29,39}, ERp44²⁹, ERp29²⁹, TMX1⁴⁰ 145

and TMX3 ²⁹, of which only ERP57, ERp72, ERp44 and ERp29 are released by platelets and
recruited to the cell surface upon activation ²⁹. The transmembrane thiol isomerase TMX1 has
been identified recently as the first thiol isomerase to negatively regulate platelet function, i.e.
inhibition of TMX1 potentiates platelets responses ⁴⁰.

Similar to ERp5 and PDI, the functions of several other surface thiol isomerases have 150 been characterized in platelets. Selective inhibition of surface ERp57 ^{38,42} also results in 151 diminished platelet activation, and thrombosis in mice. Using platelet-specific ERp57 KO 152 mice, Wang et al ⁴³ have confirmed the importance of this thiol isomerase for thrombus 153 formation in vivo. This work has also demonstrated that the C-terminal, but not the N-154 terminal, active site of ERp57 is critical for platelet aggregation. More recently, elegant work 155 using trapping mutants has identified the lectin pathway of complement activation as a novel 156 mechanism through which peri/epicellular ERp57 may regulate thromboinflammation⁴⁴. 157

ERp72 has also been shown to regulate platelet function and thrombosis in vivo ^{39,45}. 158 The role of so many similar enzymes with similar function on the platelet surface raises the 159 question as to whether different isomerases are functionally redundant. Important studies by 160 Zhou et al ³⁹ revealed that recovery of platelet aggregation of ERp72-, PDI-, and ERp57-null 161 platelets was only observed when the specific deficient thiol isomerase was reconstituted. 162 This suggests that, although similar in structure and function, thiol isomerases may act at 163 different points to sustain platelet responses and thrombosis, developing specific tasks that 164 cannot be compensated by a different isoform, i.e. thiol isomerases may work in series. It is 165 possible that there is an electron transfer chain between thiol isomerases occurring on the 166 platelet outer membrane, similar to what has been observed in the ER ⁴⁶. However, it is still 167 unclear which specific substrates are targeted by each thiol isomerase and in what sequence, 168 fostering many unanswered questions. 169

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171 2.4. Trafficking and localization of PDIs

It is presently unclear how thiol isomerases translocate to the outer membrane of cells. In eukaryotic cells, secretory proteins often follow a conventional protein secretion route, trafficking from the ER to the Golgi apparatus and subsequently to the plasma membrane ⁴⁷. However, we have shown that PDI and ERp57 do not co-localize with secretory vesicles in megakaryocyte or platelets, but are rather concentrated in a subcellular compartment near the inner surface of platelets, corresponding to the sarco/endoplasmic reticulum or dense tubular system ²⁸. Moreover, the externalization of PDI and ERp57 were highly dependent on actin polymerization, suggesting cytoskeletal rearrangement is key to the secretion of thiol isomerases in platelets ²⁸. Similar to platelets, in endothelial cells, PDI translocates to the outer membrane through Golgi-independent routes, although the precise mechanism for this has not been established ⁴⁸. In spite of these similarities, the mode of translocation may differ between cells, given that platelet thiol isomerases externalize via actin polymerization, whereas in endothelial cells actin stress fibre disruption enhanced PDI secretion ⁴⁸.

One feasible alternative yet to be investigated is the possibility that thiol isomerases 185 externalize directly through ER-plasma membrane connections ⁴⁹. In fact, these connections 186 were shown to be highly regulated by Ca^{2+} influx ⁴⁹, which is also key to signalling in 187 platelets and endothelial cells. In summary, although recent efforts have identified 188 189 mechanisms through which thiol isomerases are trafficked in cardiovascular cells, many questions remain unanswered. Understanding of how thiol isomerases are transferred to the 190 191 outer membrane of cells may enlighten, for instance, how these proteins become localized in sites of thrombosis, i.e. whether thiol isomerases are actively secreted through secretory 192 pathways or passively diffuse upon cell disruption. 193

194 **2.5.** Clinical trials with PDI inhibitors

With increasing evidence uncovering the importance of PDI to cardiovascular cells, 195 there has been growing interest to develop PDI inhibitors to treat diseases associated with 196 197 thrombosis and hypercoagulability. A comprehensive table with known inhibitors of PDI and other thiol isomerases is presented below (Table 1). In spite of this growing interest, clinical 198 trials with truly specific thiol isomerase inhibitors are currently lacking. Two phase II clinical 199 trials are underway using isoquercetin, a flavonoid that targets PDI 50 and antioxidant 200 pathways ⁵¹. The first trial aims to assess the benefits of administering isoquercetin to patients 201 with hypercoagulable states, after an initial study in healthy volunteers ⁵². The results from 202 this study (ClinicalTrials.gov Identifier: NCT01722669) have been recently published, 203 showing that daily administration of 1,000 mg isoquercetin for 56 days was able to improve 204 markers of coagulation in patients with advanced cancer ⁵⁰. Importantly, there were no 205 reports of major hemorrhages in placebo or isoquercetin-treated cohorts ⁵⁰. The second phase 206 II clinical trial using isoquercetin will explore the effects of this flavonoid in 207 thromboinflammatory biomarkers of patients with stable sickle cell 208 disease (ClinicalTrials.gov Identifier: NCT04514510). The primary outcome will measure changes in 209

the plasma soluble P-selectin levels comparing the baseline to isoquercetin response and thestudy is due to completion in October 2022.

Therefore, although significant improvements have been made to characterize PDI and other thiol isomerases in cardiovascular cells, the development of more specific inhibitors of thiol isomerases is still an ongoing drug development programme. This issue is currently being tackled with the recent discovery of novel and more selective inhibitors of PDI, as discussed below.

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218 **3. PRO-THROMBOTIC TARGETS OF THIOL ISOMERASES**

219 3.1. Integrins

220 Integrins are heterodimeric transmembrane receptors composed of an α and β subunit essential for cell migration and adhesion 53 . For instance, integrin α IIb β 3 is a platelet specific 221 receptor for fibrinogen and von Willebrand factor and is therefore essential for thrombus 222 formation. Platelet activation culminates in conformational changes in integrin aIIbB3, 223 increasing affinity for its ligands and therefore triggering thrombus formation (reviewed in 224 ¹⁰). Indeed, β 3 deficient mice have been shown to display impaired thrombosis ⁵⁴. The 225 observation that several integrins have a cysteine-rich domain, has led to the hypothesis that 226 thiol-disulphide exchange reactions may regulate the activity of these adhesion receptors. 227 228 Several thiol isomerases have been shown to associate with integrins in platelets and endothelial cells. ERp5¹⁸, ERp57⁴³ and PDI ^{54,55} have been shown to interact with integrin 229 β 3 and their activities are associated with integrin activation. 230

In contrast, two thiol isomerases, namely ERp5 and TMX1, were shown to inhibit 231 integrin β3 activation. In spite of previous reports suggesting a pro-thrombotic effect of ERp5 232 ^{18,41}, there is evidence that ERp5 catalyses the release of fibrinogen from activated integrin 233 αIIbβ3 ⁵⁶. ERp5 was shown to cleave a disulphide bond between Cys177 and Cys184 in the 234 β I domain of integrin β 3, thus leading to fibrinogen release ⁵⁶. Meanwhile, the 235 transmembrane thiol isomerase TMX1 was shown to inhibit platelet function through the 236 oxidation of integrin $\beta 3^{40}$. This reinforces the possibility that several thiol isomerases may 237 orchestrate the redox modulation of integrin β 3 on the surface of cardiovascular cells through 238 oxidizing, reducing and isomerizing cysteines on the cysteine-rich domain of integrin β 3. 239 Interestingly, the association between thiol isomerases and integrin β 3 could also be 240

perceived as bi-directional, since it has been shown that the genetic deletion of this integrin precludes PDI accumulation on the site of thrombus formation ⁵⁴. In addition, integrin β 3 has been reported to possess an endogenous reductase activity through thioredoxin-like domains ⁵⁷, similar to thiol isomerases. Therefore, thiol isomerases may regulate the activity of integrins as well integrins may regulate the exposure of thiol isomerases upon vascular injury. The precise mechanisms governing such interaction are still unclear.

Other integrins have also been reported to be mediated by PDI in vascular cells and 247 leukocytes. Lahav et al ⁵⁸ have demonstrated that inhibition with RL-90, an antibody that 248 targets PDI and to a lesser extent ERp57 ³⁸, blocked the binding of GFOGER peptide to 249 integrin a2\beta1. This provided mechanistic evidence as to how PDI and ERp57 modulate 250 adhesion of platelets to collagen surfaces, since integrin $\alpha 2\beta 1$ is an important adhesion 251 receptor for collagen ⁵⁹. Integrin regulation is not restricted to platelets only, since in 252 endothelial cells infected with dengue virus, PDI has been shown to co-localize with and 253 regulate the activation of both integrins β 1 and β 3⁶⁰. Likewise, PDI was shown to interact 254 with integrin $\alpha M\beta 2$ on the surface of neutrophils and regulate the recruitment of these cells 255 during vascular inflammation ⁶¹. 256

In spite of data showing how PDI interact and control integrins, it must be noted that 257 platelet PDI does not affect the adhesion of platelets to fibrinogen ^{37,62}. Indeed, platelet 258 adhesion to fibrinogen was shown to be mediated by GPVI ⁶³ and secondary activators, such 259 as ADP ⁶⁴. It is possible that, while PDI regulates the early activation of integrin α IIb β 3, it 260 does not affect other molecules required for sustained platelet adhesion to immobilized 261 fibrinogen, as corroborated by data of platelet-specific PDI-deficient mice ³⁷. Therefore, the 262 regulation of several integrins in cardiovascular and circulating cells exerted by PDI and 263 sibling proteins is perceived as a central mechanism through which thiol isomerases regulate 264 thrombus formation, although there are also other targets. 265

266 **3.2. GpIbα**

The adhesion receptor GpIba, part of the GPIb-IX-V complex, is the main receptor for VWF in platelets, together with the integrin α IIb β 3⁶⁵. Indeed, the relevance of GpIba has been indisputably defined since ILR4a/Gp1ba-tg mice, which lack the extracellular domain of this adhesion receptor, were shown to have impaired thrombus formation *in vivo*⁶⁵. Interestingly, the interaction between GPIba and the A1 domain of VWF was shown to be modulated by the formation of disulphide bonds in GpIba⁶⁶, suggesting that redox processes

may regulate the activation of this receptor. PDI is also capable of targeting Cys2771 and 273 Cys2773 of VWF, influencing the dimerization of VWF, which is necessary for its 274 interaction with GpIba⁶⁷. Notwithstanding, PDI was demonstrated to be in close proximity 275 with GpIba on the platelet outer membrane ⁶⁸, while the inhibition of PDI modulates the 276 exposure of free thiols in GpIba upon platelet activation, suggesting a functional association 277 between these two proteins ⁶⁸. Indeed, Stopa et al ⁶⁹ have used a kinetic-based trapping 278 approach to show that PDI interacts with GpIba, while more recently Li et al ⁷⁰ have reported 279 that PDI directly binds to GpIba on the platelet surface, catalysing the reduction of disulphide 280 bonds Cys4-Cys17 and Cys209-Cys248. This same study reported that the PDI-GPIba 281 interaction was relevant to platelet-neutrophil interaction, vascular occlusion under 282 thromboinflammatory conditions and tissue damage in ischemia-reperfusion injury. 283 Therefore, PDI has been proposed as a key regulator of GpIba on the platelet outer 284 membrane and this interaction seems relevant to various thromboinflammatory diseases. 285

286 3.3. Tissue Factor

Tissue factor (TF) is a glycoprotein key to the coagulation system, since it is the 287 cellular receptor of coagulation factors FVII and FVIIa, and the formation of the TF-FVIIa 288 complex triggers signalling events that culminate in the activation of FIX and FX 71. 289 Importantly, Cys186 and Cys209 of TF are located on the extracellular domain and capable 290 of forming a disulphide bond, which regulates the activation of TF⁷². Initial studies of 291 Ahmed et al have shown that extracellular PDI, but not ERp57, is a negative regulator of TF, 292 suppressing TF coagulant activity through a NO-dependent mechanism ⁷². This has been 293 reinforced by evidence showing that inhibition of cell-surface PDI enhances TF procoagulant 294 activity, while addition of exogenous PDI decreases TF activation in endothelial cells ⁷³. The 295 proposed mechanism of action involved the exposure of phosphatidylserine (PS), since PDI 296 addition led to increased PS exposure ⁷³, which is a known regulator of TF activation 297 (reviewed in 74). 298

However, several groups have contested the negative regulation of TF by PDI and have described opposite results. First PDI was able to enhance the procoagulant activity of TF in microvesicles in a process regulated by the chaperone activity of PDI ⁷⁵. These findings have been explored *in vivo*, showing that PDI directly promotes TF-dependent fibrin generation in a murine model of thrombosis, although the proposed mechanism involves the isomerisation of disulphide bonds in TF by reduced PDI ³². In line with these observations, PDI stimulated

the coagulant activity of TF present in extracellular vesicles secreted by endothelial cells ⁷⁵, 305 which could be related to TF-dependent fibrin deposition in vivo. More recently, Chen et al ⁷⁶ 306 have shown that PDI enhances TF-dependent thrombin generation in human peripheral blood 307 mononuclear cells, which could also propagate thrombus development upon vascular damage 308 and fibrin deposition. Indeed, it has been proposed that the two main regulators of cell-309 surface TF are PS exposure and cell-surface PDI, while there are PDI-dependent and 310 independent pathways that fine-tune the activation of TF and signalling of the coagulation 311 cascade ⁷⁴. Since thiol isomerases have a role in fibrin deposition and TF is rapidly recruited 312 to the site of intravascular damage ⁷⁷, one could speculate that the stimulation of TF by PDI 313 could represent an initial coagulation step during thrombus development. 314

Altogether, it is patent that PDI is able to interact with TF and regulate its activity, 315 316 although we still fail to fully understand how such regulation occurs. It is also unclear if other thiol isomerases facilitate TF activation, which could potentially explain contrasting results 317 obtained by different groups, since most PDI inhibitors used in previous studies were later 318 shown inhibit other thiol isomerases, e.g. PACMA31 used by Chen et al ⁷⁶ is able to inhibit 319 ERp5, ERp46, ERp57 and ERp72 78. Additional studies should be performed in platelet-320 specific PDI-deficient mice to assess if TF deposition and function are affected, or if other 321 thiol isomerases are able to overcome PDI deficiency. Likewise, it is yet unclear if platelet 322 TF (compared to other sources of TF) is relevant for thrombus development. Platelet-specific 323 double PDI/TF KO mice could also be generated to investigate if PDI and TF are 324 complementary to one another. These experiments can generate evidence to the role of the 325 PDI-TF interaction in the propagation of thrombosis and regulation of fibrin deposition in 326 vivo. 327

328 3.4. NADPH Oxidases (Noxes)

329 The Nox enzymatic system has been recently implicated as a positive regulator of platelet function ⁷⁹. This enzyme complex, which was first described in phagocytes, has been 330 identified in endothelial cells ⁸⁰, vascular smooth muscular cells (VSMC) ⁸¹ and platelets ⁸². 331 Currently, seven isoforms of Nox have been described: Nox-1, Nox-2, Nox-3, Nox-4, Nox-5, 332 333 Duox-1 and Duox-2⁸³. However, only Nox-1, Nox-2 and Nox-4 have been found in platelets ⁷⁹ although the presence of Nox4 remains a matter of debate ⁸⁴ and the potential presence of 334 other Duox proteins has not been tested. The Nox complex system consists of transmembrane 335 (gp91-phox (Nox-2) and p22-phox) as well as cytosolic subunits (p22^{phox}, p40^{phox}, p47^{phox}, 336

p67^{phox}, Noxo1, Noxa1, Rac1 and Rac2) that assemble and regulate Nox activity ⁸⁵. Upon 337 phosphorylation, the cytosolic subunits bind to the transmembrane subunit, for instance 338 p47^{phox} to p22^{phox}, through different mechanisms, depending on the subunits involved (for 339 review, see ⁸⁶). Recently, Vara et al ⁸⁷ have shown using knockout mouse models of both 340 Nox-1 and 2, that Nox-1 was the primary source of platelet superoxide downstream of GPVI, 341 whereas Nox-2 was key for responses to thrombin. Moreover, recent data suggest that 342 platelets secrete Nox-1 in platelet-derived extracellular vesicles produced upon platelet 343 activation with TRAP-6⁸⁸. 344

In this regard, PDI has been proposed as an important modulator of Nox-1 activity, 345 through a redox interaction with p47^{phox} in leukocytes ⁸⁹ and by increasing Nox-1 activation 346 in VSMC ⁹⁰ to cite two examples. In VSMC, PDI was shown to co-localize with Rac1 ⁹¹, 347 which is an essential molecule that positively regulates Nox activity (especially Nox-2) in 348 various cells ⁹². More recently, it was described that Cys400 of PDI, which is situated on the 349 C-terminal active site of PDI 93, forms a disulphide bond with Cys196 of p47^{phox} to regulate 350 Nox-1 assembly ⁹⁴. This is of particular relevance to the platelet, given that others and we 351 proposed the inhibition of the C-terminal active site of PDI as a new antithrombotic strategy 352 ^{93,95}. Indeed, we have recently shown that PDI and Nox-1 translocate to a closer proximity in 353 CRP-activated platelets and that the expression levels of these enzymes are increased in 354 platelets of individuals presenting cardiometabolic risk factors, such as obesity and high 355 blood pressure ⁹⁶. 356

Importantly, Rac1 was shown to modulate platelet hyperaggregation and endothelial dysfunction in diabetes ⁹⁷, whilst p47phox knockout mice presented limited thrombus formation ⁹⁸, indicating the Nox-regulatory proteins Rac1 and p47phox to be central in thromboinflammatory conditions. Indeed, in both studies, Nox activity was shown to be a relevant mechanism for the effects observed. Therefore it is possible that PDI may also be involved in the Rac-1-Noxes and/or p47phox-Noxes axis and alters thrombotic conditions due to the modulation of Nox activity – an alternative that is yet to be explored.

364 **3.5. Vitronectin**

³⁶⁵ Vitronectin (from Latin: *vitreous*, 'of glass') is a glycoprotein known to be relevant for ³⁶⁶ intercellular adhesion in several biological systems, including in thrombus formation ³⁶⁷ (reviewed in ⁹⁹). This glycoprotein is abundantly present in plasma (200 to 500 μ g/mL) and ³⁶⁸ within platelet α -granules ¹⁰⁰, while two different groups have shown that vitronectin-³⁶⁹ defficient mice have impaired thrombus formation *in vivo* ^{36,101}. Relevant to thrombosis,

vitronectin was shown to bind to integrins, fibrinogen, collagens, PKC, plasminogen and to 370 form a complex with thrombin and antithrombin III ⁹⁹. Importantly, there is evidence that PDI 371 catalyses the formation of the vitronectin-thrombin-antithrombin III complex in vitro 102. 372 Indeed, Bowley et al ³⁶ have shown *in vitro* that PDI can reduce disulphide bonds between 373 Cys137-Cys161 and Cys274-Cys453 of vitronectin, which would enable the binding of 374 vitronectin to integrins α IIb β 3 and α V β 3, thus sustaining thrombus formation. However, it is 375 still unclear if this process occurs on the site of vascular injury in vivo. It is also unknown if 376 vitronectin regulate thrombosis through other mechanisms, given that it is able to bind to 377 other pro-thrombotic substances, such as collagens and fibrinogen. 378

379 **3.5.** *Other targets*

The characterization of kinetic substrate-trapping techniques to identify proteins 380 capable of a physical interaction with PDI was a significant achievement in the field. In a 381 seminal paper, Stopa et al 69 demonstrated that kinetic trapping oxidized PDI variants 382 released by platelets were able to bind to GpIba, cathepsin G, glutaredoxin-1 and thioredoxin, 383 while reduced PDI variants were associated with annexin V, collagen VI, tetranectin, 384 heparanase, serpin B6, kallekrein-14 and ERp57. Through a different approach, Moretti et al 385 ¹⁰³ described an evolutionary conserved gene pairing between genes of the PDI and Rho 386 guanine-dissociation inhibitors (GDI) family of proteins. These authors have also reported a 387 physical interaction between PDI and Rho-GDI in vitro. Indeed, platelets express RhoGDI, 388 which is involved in cytoskeleton rearrangement of several eukaryotic cells, however its 389 function in thrombosis and haemostasis are still unexplored ¹⁰⁴. Finally, it was shown that 390 PDI binds to Cys374 of β -actin and that activation of integrin α IIb β 3 in the megakaryocytic 391 cell line MEG-01 was essential for the PDI-\beta-actin interaction ¹⁰⁵. Therefore, there are 392 several potential pro-thrombotic molecules that have been shown to interact with PDI. 393 Nevertheless, it remains unclear if the interaction of PDI with these molecules is relevant to 394 the process of thrombosis and haemostasis. 395

A summary of the main pro-thrombotic targets of thiol isomerases is presented inFigure 4.

398

399 4. ANTITHROMBOTIC EFFECTS OF THIOL ISOMERASES

400 In spite of substantial evidence pointing towards pro-thrombotic effects of PDI 401 ^{12,17,19,20,106,107}, this enzyme also exerts paradoxical inhibition of platelet aggregation through

NO transference in a process named transnitrosation. NO is an important platelet inhibitor 402 that acts through activating guanylate cyclase and increasing cyclic guanosine 403 monophosphate (cGMP) levels ¹⁰⁸. This induces vasodilator-stimulated phosphoprotein 404 (VASP) phosphorylation, which inactivates $\alpha IIb\beta 3$ ¹⁰⁹⁻¹¹¹. The discovery that PDI has 405 denitrosation activity was first reported by the Mutus laboratory that showed that S-406 nitrosothiols (RSNOs) inhibit platelets through a dual mechanism: first through denitrosation 407 of RSNOs by PDI, thus releasing NO and secondly due to a direct RSNO reaction with PDI, 408 rendering it unable to perform disulphide exchange on the platelet membrane ^{112,113}. Indeed it 409 was recently demonstrated in a cell-free environment that 57% of peroxynitrite, which is the 410 product of the reaction between NO and superoxide, oxidizes PDI through a 2-electron 411 mechanism while 43% is converted to nitrate and other radicals ¹¹⁴. This was further studied 412 by other groups that showed that different NO donors attenuate platelet function through 413 PDI-mediated denitrosation ^{115,116}. More recently, Bekendam et al have proposed that the S-414 nitrosylation of vascular thiol isomerases PDI, ERp5 and ERp57 by NO is able to at least 415 partially mediate vascular quiescence through the inhibition of these thiol isomerases ¹¹⁷. 416 417 Altogether, the pro-thrombotic activity of PDI seems to overcome its inhibitory effect in physiological scenarios – whether this would hold true in the context of disease is yet to be 418 419 defined.

In addition to S-transnitrosation of thiol isomerases, it has been recently described that 420 some thiol isomerases can exert a negative regulation of platelet function, i.e. inhibit platelet 421 activation. The only thiol isomerase protein described to exert such inhibitory effect thus far 422 is the transmembrane TMX1. This protein was first detected in megakaryocytes over 10 years 423 ago, together with other isoforms, namely: TMX2, TMX3 and TMX4¹². TMX1 helps with 424 protein folding in the ER through a CPAC-active site (in contrast to the CGHC active site of 425 PDI, ERp5, ERp57 and ERp72) through the formation of disulphide bonds in newly formed 426 proteins ¹¹⁸. Elegant work performed by Zhao et al ⁴⁰ using TMX1-deficient platelets as well 427 as recombinant TMX1 addition have demonstrated that TMX1 decreases platelet and 428 thrombotic responses through the oxidation of integrin a_IIbβ3. Moreover, addition of an anti-429 TMX1 antibody potentiated platelet aggregation, while addition of recombinant TMX1 430 inhibited platelet aggregation exerted by different agonists ⁴⁰. Therefore, it is possible that 431 other transmembrane thiol isomerases found on the platelet membrane are also able to 432 negatively regulate platelet responses. The investigation of the effects and possible inter-433 regulation of different thiol isomerases present on the platelet outer surface will allow for a 434 more comprehensive understanding of how this family of proteins may modulate 435

thromboinflammatory conditions. In this regard, the discovery of selective inhibitors of thiolisomerases will greatly advance the field.

438

439 **5. INHIBITORS OF THIOL ISOMERASES**

440 **5.1** *Small molecule inhibitors*

Considering the deleterious effects of PDI and other thiol isomerases to the 441 cardiovascular system ^{14,16,36,42,44}, there has been great scientific effort to identify novel, non-442 443 toxic and selective inhibitors of thiol isomerases. Indeed, many compounds have been identified over the last decades (an up to date summary of PDI inhibitors is presented in 444 Table 1). These small molecule inhibitors were often characterized through high throughput 445 screening of chemical libraries. Frequently, PDI inhibitors were tentatively identified from 446 compounds screened for neurodegenerative diseases or cancer. For instance, Hoffstrom et al 447 ¹¹⁹ screened 68,887 compounds against a cell based model for Huntington's disease and 448 found that PDI was the molecular target for the top 5 hits, which included 16F16, a 449 compound that was later shown to bind covalently to Cys36 and Cys39 of the N-terminal 450 active site of PDI 120. LOC14, a reversible inhibitor of PDI, was also identified after 451 screening for potent rescue of a Huntington's disease cell based model ¹²⁰. In a similar 452 approach, Vatolin et al ¹²¹ screened 30,335 compounds for activity against *in vivo* and *in vitro* 453 models of multiple myeloma and have identified CCF642 as a lead compound. This same 454 report used the di-eosin reductase assay to show that CCF642 was able to inhibit PDI and 455 other thiol isomerases at low micromolar concentrations, although a recent report has shown 456 a much higher IC₅₀ for CCF642 in the insulin turbidimetry assay against several thiol 457 isomerases ⁷⁸. It is possible that discrepant results using CCF642 were due to different assays 458 being employed to characterize the anti-PDI activity. This notion is corroborated by 459 Bekendam et al ¹²² who have elegantly shown that reversible PDI inhibitors bepristat 1a and 460 bepristat 2a, which were identified after a high-throughput screening of 348,505 compounds, 461 were able to inhibit PDI activity only when this was assessed using the insulin turbidimetry 462 463 assay.

In parallel to PDI inhibitors identified after screening for neurodegenerative or cancer diseases, several groups have conducted high throughput screening in which the primary screen consisted of PDI reductase assay. This is the case for AS15, an aminobenzylphenol compound which covalently binds and inhibits PDI at nanomolar concentrations and

decreases cell proliferation of Glioblastoma cell lines ¹²³. In a similar approach, this same 468 group has identified 35G8, which is another nanomolar inhibitor of PDI that also inhibits 469 proliferation of Glioblastoma cell lines ¹²⁴. Importantly, 35G8 was shown to covalently bind 470 to Cys397 of the C-terminal active site of PDI ¹²⁴, which has been shown by us ¹²⁵ and others 471 ¹²⁶ to be a relevant target site to limit the pro-thrombotic actions of PDI in platelets. In 472 contrast, KSC-34 has been described as the only inhibitor to be ~30 times more selective 473 towards the N-terminal over the C-terminal active site of PDI ¹²⁷. One could hypothesize that 474 different inhibitors that target different parts of PDI could exert opposing effects in 475 thrombosis and haemostasis, however at present neither KSC-34 nor 35G8 have been tested 476 in platelets or other cardiovascular cells. Indeed, the majority of small molecule inhibitors 477 described thus far have not been tested for their effects in thrombosis and haemostasis. These 478 include: 16F16¹¹⁹, 35G8¹²⁴, AS15¹²³, BAP1 and BAP2¹²⁸, CCF642¹²¹, E64FC26⁷⁸, KSC-479 34¹²⁹, LOC14¹²⁰, Origamicin¹³⁰, Securinine¹³¹, SK053¹³² and STK076545¹³³. Recent data 480 from our lab suggests that LOC14 exerts anti-platelet effects, while CCF642 and 16F16 do 481 not alter platelet function (data not shown). It would be important to assess the anti-platelet 482 483 potential of other PDI inhibitors.

Only five small molecule inhibitors have been assessed for their effects in thrombosis 484 and haemostasis. Bepristats are selective and reversible inhibitors of PDI with an IC₅₀ ranging 485 from 0.7 to 1.2 µM against the reductase activity of PDI measured through insulin 486 turbidimetry ¹²², while PACMA-31 is an irreversible micromolar inhibitor of PDI ^{122,134}. In 487 spite of the low IC₅₀ for PDI reductase activity, Bekendam et al have shown that 30 μ M of 488 bepristats or PACMA-31 was able to abrogate platelet aggregation, while bepristats exerted 489 no effect in P-selectin exposure ¹²². Unpublished data from our lab suggest that 490 concentrations as low as 7.5 µM of bepristat 2a can inhibit platelet aggregation, activation 491 and calcium mobilization, depending on the agonist used. Such discrepancy between the 492 concentration needed to inhibit PDI in a cell-free system and the one needed to inhibit 493 platelets was also found for ML359. This inhibitor was able to marginally inhibit thrombin-494 induced platelet aggregation (25%) at 30 μ M ¹³⁵ in spite of an IC₅₀ over 100 times lower for 495 PDI reductase activity, suggesting poor biochemical properties or off-target effects ¹³⁶. On the 496 other hand, HPW-RX40 has shown similar low micromolar IC₅₀ for both PDI reductase assay 497 measured through the di-eosin assay and platelet aggregation induced by several agonists ¹³⁷. 498 Finally, we have recently shown that the cysteinyl LT receptor antagonist zafirlukast is a pan 499 inhibitor of thiol isomerases, decreasing the reductase activity of PDI, ERp5, ERp57, ERp72 500 and TRX at micromolar concentrations ¹³⁸. Similar concentrations of zafirlukast were able to 501

inhibit platelet aggregation, activation, calcium mobilization and *in vivo* thrombosis with no
 effect on bleeding time ¹³⁸. Therefore, there are currently few small molecule inhibitors of
 PDI and other thiol isomerases with well-described effects in thrombosis and haemostasis.

In addition to the five small molecule inhibitors that have been tested in platelets, 505 bacitracin was initially perceived as a selective PDI inhibitor. However, this was challenged 506 over 10 years ago, when Karala and Ruddock definitively showed that bacitracin is not a 507 selective inhibitor of PDI, neither does it exert its cellular effects through the inhibition of 508 thiol isomerases ¹³⁹. Therefore, the identification of novel inhibitors of PDI faces several 509 challenges. First, it is possible that current molecules perceived as PDI inhibitors do not exert 510 their anti-platelet effect through targeting this enzyme, similar to bacitracin. This is 511 corroborated by literature exposed above that show that bepristats and PACMA-31 need a 512 much higher concentration to inhibit platelets than to inhibit thiol isomerases in cell-free 513 environments ¹²². Definitive proof of the specificity of inhibitors could be achieved through 514 515 the use of platelet PDI-deficient models. Secondly, it is yet unknown if small molecule inhibitors described as anti-cancer agents have effects on thrombosis and haemostasis. Lastly, 516 although there are a few PDI-selective inhibitors, such as bepristats, there are no selective 517 inhibitors for other thiol isomerases. The identification of such compounds would forward the 518 field as it has been shown that different thiol isomerases may have distinct modes of action in 519 platelets ³⁹. In conclusion, it would be beneficial if future studies: 1) prove that current and 520 future PDI inhibitors act through targeting a specific thiol isomerase, 2) investigate if the 521 anti-cancer and anti-platelet properties of thiol isomerase inhibitors overlap and 3) identify 522 inhibitors that target specific protein activities (reductase, oxidase, isomerase and chaperone 523 activity). 524

525 5.2 Flavonoids and natural compounds

Similar to small molecule inhibitors, flavonoids and other natural compounds have 526 been extensively studied as potential inhibitors of thiol isomerases. The most prominent and 527 clinically advanced natural compound is isoquercetin, which is a derivative of the flavonoid 528 quercetin currently being employed in phase II clinical trial as a potential anti-thrombotic 529 drug as described above ⁵⁰. However, rutin, which is a quercetin derivative containing a 530 rutinose lateral chain in carbon 3, was the first derivative proposed as a promising PDI 531 inhibitor ¹⁴⁰. Jasuja et al ¹⁴⁰ demonstrated the ability of rutin to inhibit thrombosis in mice at 532 concentrations as low as 0.1 mg/kg. This same group have reported that rutin binds reversibly 533

to the b'x domain of PDI, similar to bepristats ^{122,141}. Recently, we have reported that myricetin, which is a flavonoid of similar structure to quercetin, is also able to inhibit platelets at low micromolar concentrations ⁶². This was attributed to the inhibition of PDI and ERp5, since this flavonoid was shown to inhibit these enzymes at similar concentrations needed to achieve platelet inhibition ⁶², although definitive proof is still needed.

Other natural compounds have also been proposed as novel inhibitors of thiol 539 isomerases. For instance, rosmarinic acid, commonly found in Danshen (Salva miltiorrhiza) 540 was shown to inhibit ERp57 and promote platelet inhibition at low micromolar 541 concentrations, depending on the agonist used ¹⁴². Punicalagin was also shown to inhibit 542 ERp57 at low micromolar concentrations in a cell-free environment, while the biological 543 actions of this polyphenolic compound were lost in ERp57-silenced neuroblatoma cells, 544 545 suggesting ERp57 to be the main target of punicalagin in cellular systems ¹⁴³. Similarly, tannic acid was demonstrated to bind to PDI with high affinity, after a directed in silico 546 screening of over 60 natural compounds ¹⁴⁴. This study demonstrated that tannic acid inhibits 547 several thiol isomerases and prevents thrombus formation in the cremaster laser-induced 548 model of thrombosis *in vivo*¹⁴⁴. Therefore, several natural compounds have been identified as 549 potent inhibitors of PDI and other thiol isomerases, with implications to thrombosis and 550 haemostasis. However, it is still unclear if thiol isomerase inhibition is indeed the mechanism 551 of action of these compounds in biological systems. 552

Juglone, which is an allelopathic compound present in the roots of walnut trees, has 553 been shown to inhibit platelet aggregation, possibly through the inhibition of both PDI and 554 Akt ¹⁴⁵. Indeed, other flavonoids, such as quercetin, apigenin and catechin have been shown 555 to act as kinase inhibitors and to inhibit the activity of Src family kinases in platelets ¹⁴⁶. A 556 previous study of our lab showed that quercetin and other structurally related flavonoids were 557 able to interact with fibrinogen and collagen, to prevent Syk phosphorylation and to be 558 internalized by megakaryocytes and platelets ¹⁴⁷. Likewise, quercetin, catechin and other 559 structurally related flavonoids were shown to inhibit platelet aggregation, and act as 560 competitive of the thromboxane A₂ (TxA₂) receptor ¹⁴⁸. Therefore, similar to small molecule 561 inhibitors, it is still unclear if the biological activity of natural compounds described as PDI 562 inhibitors is indeed due to thiol isomerase inhibition. A thorough analysis of which 563 compounds exert their effect through thiol isomerase inhibition and which thiol isomerases 564 are involved would greatly benefit the development of more effective compounds to treat and 565 prevent thrombosis. 566

567 5.3 Peptide inhibitors

Peptides have been used to treat diseases for nearly 100 years, since insulin was first 568 isolated and commercialized (for review, see ¹⁴⁹). However, there is currently only one 569 peptide inhibitor described to inhibit thiol isomerases. The peptide CxxCpep was first 570 synthesized by de A. Paes et al ¹⁵⁰ as the 12 amino-acid sequence of the CGHC-redox active 571 site of PDI (VEFYAPWCGHCK). These authors have shown that CxxCpep was able to 572 inhibit PDI in neutrophils, thus decreasing the assembly of NADPH oxidase complexes ¹⁵⁰. 573 We have expanded these studies and shown that CxxCpep inhibits platelets and binds to 574 Cys397 and Cys400 of the C-terminal active site 125 – supporting the notion that the pro-575 thrombotic effects of PDI are orchestrated by the C-terminal redox active site ^{37,126}. In 576 addition, we have evidence that this peptide is membrane impermeable (unpublished), 577 578 reiterating that the extracellular pool of thiol isomerases is important to regulate platelet function. However, definitive proof is still lacking to ascertain the specificity of CxxCpep 579 580 towards PDI. A summary of PDI inhibitors with known binding sites is presented in Figure 5. Future research should design peptides to selectively inhibit other thiol isomerases. These 581 inhibitors could serve as templates for the development of stable, selective and non-toxic 582 peptide inhibitors. 583

584

585 6. FUTURE PERSPECTIVES

There has been great scientific advancement since PDI was first identified in 1963 ²¹ 586 and its protein sequence determined in 1985²⁵. Several decades later, it is now undisputed 587 that PDI and other thiol isomerases control platelet function, acting as an orchestrating force 588 in the complex and dynamic process of thrombosis and haemostasis. In parallel, there has 589 been a surge of novel inhibitors of thiol isomerases discovered through high-throughput 590 screening of small molecules, such as bepristasts ¹²² and repurposing of drugs currently used 591 in other settings, such as zafirlukast ¹³⁸. However, there are still pressing questions left 592 unanswered in order to translate basic findings to the clinic. 593

594 First, it would be important to understand which molecules are targeted by thiol 595 isomerases and how these interactions occur. It is widely accepted that PDI, ERp5, ERp57 596 and ERp72 regulate integrin β 3 activation, while TMX1 acts as a negative regulator, as 597 exposed above in **subheading 3.1**. Interestingly, platelet aggregation in ERp72-, PDI-, and 598 ERp57-null platelets was only recovered when the deleted thiol isomerase was added back ³⁹, 599 suggesting that each enzyme acts in series, targeting different molecules or different parts of 600 the same molecules. This also reinforces the notion that thiol isomerases are not redundant in 601 platelets. Another feasible alternative is that thiol isomerases interact amongst themselves on 602 the outer surface of cardiovascular cells. Moreover, it is unclear if ERp44, ERp29 and TMX3, 603 which were found in platelets ²⁹, are also able to influence thrombosis and haemostasis.

Second, novel research should address the possible overlap between the anticancer, 604 neuroprotective and antiplatelet properties of thiol isomerase inhibitors. Specifically, it would 605 be interesting to know if inhibitors that target different parts of PDI are able to modulate 606 different aspects regulated by PDI. For instance, the C-terminal, but not the N-terminal, 607 active site of PDI is required to modulate platelets ^{17,93}. In parallel, the neuroprotective 608 inhibitor 16F16 targets the N-terminal active site ¹²⁰, while the anticancer inhibitor 35G8 609 targets the C-terminal active site of PDI ¹²⁴, therefore, it would be important to understand 610 how these and other inhibitors regulate platelet function. Such characterization would deepen 611 612 our knowledge on possible side effects of thiol isomerase inhibitors and propose a template for the development of more selective compounds. 613

Finally, the characterization of selective inhibitors for each thiol isomerase and 614 translation of these inhibitors to the clinic are of great interest to the field. Currently there are 615 a few inhibitors that are selective to PDI over sibling proteins, such as 16F16 ^{119,120} and 616 bepristats ¹²², however there is no such equivalent to ERp5, ERp57 or ERp72. In addition, 617 full characterization of off-target effects of these inhibitors in vivo is still lacking. It is also 618 unclear how thiol isomerases in platelets are correlated to thrombosis in pro-thrombotic 619 conditions, such as metabolic syndrome and cancer. Promising findings of Zwicker et al ⁵⁰ 620 621 have shown a potential benefit of using isoquercetin to prevent cancer-associated thrombosis and it is expected that this positive outcome will bring interest to the development of 622 selective inhibitors of other thiol isomerases. 623

In conclusion, thiol isomerases are central to many biological systems and could be perceived as a driving force that dictates thrombus development. This complex interplay involves redox reactions with key adhesion receptors occurring at the platelet outer membrane. Ultimately, it becomes increasingly evident that platelets are highly regulated by redox processes, while novel techniques, inhibitors and other tools are fostering exciting discoveries in this rapid-growing field.

631 AUTHOR CONTRIBUTIONS

R.S. Gaspar wrote the manuscript and drafted the figures, while J.M. Gibbins wrote andrevised the manuscript. All authors approved the final version submitted.

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638 were made using BioRender.com.



Figure 1. Inside-out and outside-in signalling in platelets. Platelet activation can be 642 didactically divided in two pathways: inside-out and outside-in signalling. Inside-out 643 signalling refers to binding of agonists to their respective receptors on the platelet membrane 644 (e.g. collagen binding to GPVI). This initial binding will lead to specific pathways of each 645 receptor that will culminate in a common pathway that involves increased intracellular Ca²⁺ 646 mobilisation, granule secretion, activation of protein kinase C (PKC), phosphoinositide 3-647 kinase (PI3K) and mitogen-activated protein kinases (MAPKs). These molecules and 648 signalling events will then activate integrin aIIb₃ in a process that requires protein 649 disulphide isomerase (PDI) as well as other thiol isomerases. Upon binding to fibrinogen, the 650 integrin aIIb_{β3} will cause a series of intracellular signalling events, termed outside-in 651 signalling, that will potentiate initial response by agonists. PAR: protease-activated receptor. 652 ADP: adenosine diphosphate. TP: thromboxane receptor. TxA₂: thromboxane A₂. 653



Figure 2. Protein disulphide isomerase catalyses redox reactions. The 3D structures of 656 protein disulphide isomerase (PDI) were obtained from the PDB database (PDB ID: 4EL1 for 657 oxidized and PDB ID: 4EKZ for reduced). PDI has 4 domains and an x-linker to promote 658 flexibility. Its catalytic sites are located in a and a'-domains. Due to its particular structure, 659 PDI can catalyse the reduction of disulphide bonds into free thiols, oxidation of thiols into 660 disulphide bonds or isomerisation of disulphide bonds, leading to a different protein 661 conformation. Alternatively, it can also transfer nitric oxide (NO) from nitrosothiols (SNO) 662 between proteins in a process named transnitrosation. 663



665

Figure 3. Historical landmarks of the PDI field. Decades of intensive research output led to many discoveries linking PDI and sibling proteins to thrombosis and haemostasis. From its initial characterisation in the 1960s-1980s, PDI was later found on the outer surface of cardiovascular cells. Other thiol isomerases were also identified and collectively shown to regulate thrombosis. More recently, we started to uncover how these proteins become externalized while a feasible candidate for drug development is currently being tested in clinical trials.



Figure 4. Pro-thrombotic targets of thiol isomerases. PDI, ERp5, ERp57 and ERp72 are 675 known regulators of platelet function through their interaction with integrin aIIb₃. In 676 contrast, TMX1 has been shown to inhibit integrin aIIb_{β3}, being the first thiol isomerase 677 described to negatively regulate thrombosis. PDI is also able to interact with integrin $\alpha 2\beta 1$, 678 Von Willebrand Factor (VWF), GpIba, Nox-1, Nox-2, tissue factor and vitronectin all of 679 which are pro-thrombotic. Therefore, the interaction of thiol isomerases with these pro-680 thrombotic molecules is a feasible mechanism through which thiol isomerases can control 681 platelet function. Other proteins, such as β-actin and RhoGDI were also shown to interact 682 with PDI, however the relevance of this interaction to thrombosis and haemostasis is still 683 unclear. It is also unclear if there are other targets that could contribute to the pro-thrombotic 684 (or anti-thrombotic for TMX1) effect of thiol isomerase proteins. 685



Figure 5. PDI inhibitors with known binding sites. Several PDI inhibitors have been characterized, however only a few of these have a known binding site.16F16, KSC-34 and LOC14 bind close to the a'-active site; CxxCpep, 35G8 and PACMA-31 bind close to the aactive site, while rutin, bepristats, BAPs and isoquercetin bind near the b' and x active sites. There are currently no inhibitors that selectively target the b-domain.

694	Table 1. List of known thiol isomerase inhibitors and their effects on thrombosis and
695	haemostasis.

Inhibitor	TI targeted	IC ₅₀ for TI inhibition	Binding site	Effects on thrombosis and haemostasis	ref
Small					
molecule					
inhibitors					
16F16	PDI ERp5 ERp57 ERp72	6 μM for PDI 50 μM for ERp72 30 μM for ERp5	Cys36 and Cys39 of the N-terminal active site of PDI	Platelet function - *	120 119
35G8	PDI	0.17 μM for PDI	Cys397 of the C-terminal active site of PDI		124
AS15	PDI	0.3 µM	Not tested	Not tested	123
BAP1 BAP2	PDI	0.83 μM 0.93 μM	His256 b'domain	Not tested	128
Bepristat 1a Bepristat 2a	PDI	0.7 μM 1.2 μM	b'x domain	Platelet function <i>In vivo</i> thrombosis	122
		$100 \ \mu M$ for	Near C-		
CCF642	PDI	PDI**	terminal	Platelet	121
	ERp5	100 μM for ERp5**	active site of PDI [#]	function - *	78
	PDI	$2 \ \mu M$ for PDI			
E64FC26	ERp5	$25 \ \mu M$ for	Not tested	Not tested	78
	ERp57	ERp5			

	ERp72	20 μM for ERp57 25 μM for ERp72			
HPW-RX40	PDI ERp5 ERp57 ERp72	1.45 μM for PDI 2.6 μM for ERp5 4.3 μM for ERp57 18.8 μM for ERp72	Near C- terminal active site of PDI [#]	Platelet function <i>In vitro</i> thrombosis	137
KSC-34	PDI	3.5 μM for PDI	Cys53 of the N-terminal active site of PDI	Not tested	129
LOC14	PDI ERp5 ERp57 ERp72	150 μM for PDI 45 μM for ERp5 4.97 μM for ERp57 100 μM for ERp72	Near the N- terminal active site of PDI	Platelet function *	120 151 78
ML359	PDI	0.25 μM for PDI	Not tested	Platelet function	135
Origamicin	PDI ERp5 ERp57 ERp72	Not tested	Not tested	Not tested	130
PACMA31	PDI	$7 \ \mu M$ for PDI	Cys397 and	Platelet	78

	ERp5	5 µM for	Cys400 of the	function	134
	ERp46	ERp5	C-terminal	In vitro	152
	ERp57	$20 \ \mu M$ for	active site of	thrombosis	122
	ERp72	ERp72	PDI	Haemostasis	
		Not tested for			
		ERp57 and			
		ERp46			
			Near the N-		
Saauvinina	ורום	Not tostad	terminal	Not tostad	131
Securimine	PDI	Not tested	active site of	Not lested	
			PDI [#]		
			Near the C-		
SK053	ורום	$10 \ \mu M$ for	terminal	Not tostad	132
5K055	FDI	PDI	active site of	Not lested	
			PDI [#]		
		2.16 µM for			
STK076545	PDI	PDI	Not tested	Not tested	133
	PDI			Platelet	
	ERp5			function	
Zafirlukast	ERp57	Not tested	Not tested	In vivo	138
	ERp72			thrombosis	
	TRX			Haemostasis -	
Flavonoids					
and natural					
compounds					
	PDI			Platelet	
ADTM	ERp5	Not tested	Not tested	function	153
	ERp57	110000000	1.00.00000	In vivo	
	ERp72			thrombosis	
Gallovlated			Near the	Platelet	
catechins	ERp57	Not tested	active sites of	function	154,155
cure chillip			ERp57 [#]		

				Platelet		
Ingland	PDI	1.61 µM	Not to stad	function	145	
Jugione			Not tested	In vitro	110	
				thrombosis		
		0.15 μM for PDI				
Juniferdin	PDI ERp5	5 uM for		Platelet	156	
epoxide		ERp5	Not tested	function	135	
I	Thioredoxin	3 M for				
		Thioredoxin				
			NI 41	Platelet		
	זרום	Not tested	Near the	function	62	
Myricetin	ERp5		PDI and	In vitro		
			ERn5 [#]	thrombosis		
			Liqu	Haemostasis -		
Punicalagin	ERp57	1 µM	Not tested	Not tested	143	
Rosmarinic	ERp57	176 μΜ	Near the	Platelet	142	
acid			active sites of	function		
			ERp57#	Distalat		
				function		
Rutin	PDI 7-	7-10 µM	h'x domain	In vivo	78,140,141,157	
Kutin			o A domum	thrombosis		
				Haemostasis -		
				Platelet		
	PDI	Not tested	Near the C-	function		
Tannic acid			terminal	In vivo	144	
			active site of	thrombosis		
			PDI"	Haemostasis -		
Peptide						
inhibitors						
СххСрер	PDI	Not tested	Cys400 of	Platelet	125	
			reduced PDI	function		

	Mastanavan	ורות	Not tosto	Not tostad		Platelet		158,159	
	Mastoparan	FDI	Not lester	1	not tested		function		
696	TI: Thiol ison	nerase. *Prelin	minary data	from	our	lab.	**:	Disputed.	:Decreased.
697	:Increased: Unaffected. #: Predicted.								

699 **REFERENCES**

- Bizzozero, J. Ueber einen neuen Formbestandtheil des Blutes und dessen Rolle bei
 der Thrombose und der Blutgerinnung. Archiv für pathologische Anatomie und
 Physiologie und für klinische Medicin 90, 261-332 (1882).
- 2 Quick, A. J. *Bleeding problems in clinical medicine*. (Saunders, 1970).
- Roskam, J., Hugues, J. & Bounameaux, Y. LHEMOSTASE SPONTANEE ETUDE
 SYNTHETIQUE ET ANALYTIQUE. JOURNAL DE PHYSIOLOGIE 53, 175-&
 (1961).
- Bettex-Galland, M. & Lüscher, E. Thrombosthenin—a contractile protein from thrombocytes. Its extraction from human blood platelets and some of its properties.
 Biochimica et biophysica acta 49, 536-547 (1961).
- Zahn, F. W. Untersuchungen über thrombose. Archiv für pathologische Anatomie und
 Physiologie und für klinische Medicin 62, 81-124 (1874).
- de Gaetano, G. Historical overview of the role of platelets in hemostasis and
 thrombosis. *Haematologica* 86, 349-356 (2001).
- 714 7 Bye, A. P., Unsworth, A. J. & Gibbins, J. M. Platelet signaling: a complex interplay
 715 between inhibitory and activatory networks. *Journal of Thrombosis and Haemostasis*716 14, 918-930 (2016).
- Savage, B., Saldivar, E. & Ruggeri, Z. M. Initiation of platelet adhesion by arrest onto
 fibrinogen or translocation on von Willebrand factor. *Cell* 84, 289-297 (1996).
- Massberg, S. *et al.* A crucial role of glycoprotein VI for platelet recruitment to the
 injured arterial wall in vivo. *J Exp Med* 197, 41-49 (2003).
- Durrant, T. N., van den Bosch, M. T. & Hers, I. Integrin αIIbβ3 outside-in signaling.
 Blood 130, 1607-1619 (2017).
- Mancuso, M. E. & Santagostino, E. Platelets: much more than bricks in a breached
 wall. *Br J Haematol* 178, 209-219, doi:10.1111/bjh.14653 (2017).
- Holbrook, L. M. *et al.* Platelets release novel thiol isomerase enzymes which are
 recruited to the cell surface following activation. *Br J Haematol* 148, 627-637,
 doi:10.1111/j.1365-2141.2009.07994.x (2010).
- Jasuja, R., Furie, B. & Furie, B. C. Endothelium-derived but not platelet-derived protein disulfide isomerase is required for thrombus formation in vivo. *Blood, The Journal of the American Society of Hematology* 116, 4665-4674 (2010).
- 731 14 Chen, K., Lin, Y. & Detwiler, T. C. Protein disulfide isomerase activity is released by
 732 activated platelets. (1992).
- Wu, Y. & Essex, D. W. Vascular thiol isomerases in thrombosis: The Yin and Yang. *Journal of Thrombosis and Haemostasis* (2020).
- Gaspar, R. S., Trostchansky, A. & Paes, A. M. Potential Role of Protein Disulfide
 Isomerase in Metabolic Syndrome-Derived Platelet Hyperactivity. *Oxid Med Cell Longev* 2016, 2423547, doi:10.1155/2016/2423547 (2016).
- Kim, K. *et al.* Platelet protein disulfide isomerase is required for thrombus formation
 but not for hemostasis in mice. *Blood* 122, 1052-1061, doi:10.1182/blood-2013-03492504 (2013).
- 741 18 Jordan, P. A. *et al.* A role for the thiol isomerase protein ERP5 in platelet function.
 742 *Blood* 105, 1500-1507, doi:10.1182/blood-2004-02-0608 (2005).
- Holbrook, L. M. *et al.* The platelet-surface thiol isomerase enzyme ERp57 modulates
 platelet function. *J Thromb Haemost* 10, 278-288, doi:10.1111/j.15387836.2011.04593.x (2012).
- Holbrook, L. M. *et al.* A humanized monoclonal antibody that inhibits platelet-surface
 ERp72 reveals a role for ERp72 in thrombosis. *J Thromb Haemost* 16, 367-377,
 doi:10.1111/jth.13878 (2018).

- Venetianer, P. & Straub, F. The enzymic reactivation of reduced ribonuclease. *Biochimica et Biophysica Acta (BBA)-Specialized Section on Enzymological Subjects*67, 166-168 (1963).
- Goldberger, R. F., Epstein, C. J. & Anfinsen, C. B. Acceleration of reactivation of reduced bovine pancreatic ribonuclease by a microsomal system from rat liver. *Journal of biological chemistry* 238, 628-635 (1963).
- Hawkins, H. C. & Freedman, R. B. Randomly reoxidised soybean trypsin inhibitor
 and the possibility of conformational barriers to disulphide isomerization in proteins. *FEBS letters* 58, 7-11 (1975).
- Tomizawa, H. H. Mode of action of an insulin-degrading enzyme from beef liver.
 Journal of Biological Chemistry 237, 428-431 (1962).
- Edman, J. C., Ellis, L., Blacher, R. W., Roth, R. A. & Rutter, W. J. Sequence of
 protein disulphide isomerase and implications of its relationship to thioredoxin. *Nature* 317, 267-270 (1985).
- Essex, D. W., Chen, K. & Swiatkowska, M. Localization of protein disulfide
 isomerase to the external surface of the platelet plasma membrane. (1995).
- Manickam, N., Sun, X., Li, M., Gazitt, Y. & Essex, D. W. Protein disulphide
 isomerase in platelet function. *British journal of haematology* 140, 223-229 (2008).
- Crescente, M. *et al.* Intracellular trafficking, localization, and mobilization of platelet borne thiol isomerases. *Arteriosclerosis, thrombosis, and vascular biology* 36, 1164 1173 (2016).
- Holbrook, L. M. *et al.* Platelets release novel thiol isomerase enzymes which are
 recruited to the cell surface following activation. *British journal of haematology* 148,
 627-637 (2010).
- Raturi, A., Miersch, S., Hudson, J. W. & Mutus, B. Platelet microparticle-associated
 protein disulfide isomerase promotes platelet aggregation and inactivates insulin. *Biochimica et Biophysica Acta (BBA)-Biomembranes* 1778, 2790-2796 (2008).
- Hotchkiss, K. A., Matthias, L. J. & Hogg, P. J. Exposure of the cryptic Arg-Gly-Asp
 sequence in thrombospondin-1 by protein disulfide isomerase. *Biochimica et Biophysica Acta (BBA)-Protein Structure and Molecular Enzymology* 1388, 478-488
 (1998).
- Reinhardt, C. *et al.* Protein disulfide isomerase acts as an injury response signal that
 enhances fibrin generation via tissue factor activation. *The Journal of clinical investigation* 118, 1110-1122 (2008).
- Tanaka, L. Y. *et al.* Peri/epicellular protein disulfide isomerase-A1 acts as an
 upstream organizer of cytoskeletal mechanoadaptation in vascular smooth muscle *American Journal of Physiology-Heart and Circulatory Physiology* 316, H566H579 (2019).
- Tanaka, L. Y. *et al.* Peri/epicellular protein disulfide isomerase sustains vascular
 lumen caliber through an anticonstrictive remodeling effect. *Hypertension* 67, 613622 (2016).
- Cho, J., Furie, B. C., Coughlin, S. R. & Furie, B. A critical role for extracellular
 protein disulfide isomerase during thrombus formation in mice. *The Journal of clinical investigation* 118, 1123-1131 (2008).
- Bowley, S. R., Fang, C., Merrill-Skoloff, G., Furie, B. C. & Furie, B. Protein disulfide
 isomerase secretion following vascular injury initiates a regulatory pathway for
 thrombus formation. *Nature communications* 8, 1-13 (2017).
- Kim, K. *et al.* Platelet protein disulfide isomerase is required for thrombus formation
 but not for hemostasis in mice. *Blood, The Journal of the American Society of Hematology* 122, 1052-1061 (2013).

- Wu, Y. *et al.* The disulfide isomerase ERp57 mediates platelet aggregation,
 hemostasis, and thrombosis. *Blood, The Journal of the American Society of Hematology* 119, 1737-1746 (2012).
- 39 Zhou, J. *et al.* The disulfide isomerase ERp72 supports arterial thrombosis in mice.
 803 *Blood* 130, 817-828 (2017).
- 80440Zhao, Z. et al. The transmembrane protein disulfide isomerase TMX1 negatively805regulates platelet responses. blood 133, 246-251 (2019).
- Passam, F. H. *et al.* Both platelet-and endothelial cell-derived ERp5 support thrombus formation in a laser-induced mouse model of thrombosis. *Blood, The Journal of the American Society of Hematology* 125, 2276-2285 (2015).
- HOLBROOK, L. M. *et al.* The platelet-surface thiol isomerase enzyme ERp57
 modulates platelet function. *Journal of Thrombosis and Haemostasis* 10, 278-288
 (2012).
- Wang, L. *et al.* Platelet-derived ERp57 mediates platelet incorporation into a growing
 thrombus by regulation of the αIIbβ3 integrin. *Blood* 122, 3642-3650 (2013).
- Eriksson, O. *et al.* Thiol isomerase ERp57 targets and modulates the lectin pathway of
 complement activation. *Journal of Biological Chemistry* 294, 4878-4888 (2019).
- 45 Holbrook, L. M. *et al.* A humanized monoclonal antibody that inhibits platelet-surface
 817 ER p72 reveals a role for ER p72 in thrombosis. *Journal of Thrombosis and*818 *Haemostasis* 16, 367-377 (2018).
- 46 Araki, K. *et al.* Ero1-α and PDIs constitute a hierarchical electron transfer network of
 820 endoplasmic reticulum oxidoreductases. *Journal of Cell Biology* 202, 861-874 (2013).
 821 47 Viotti, C. in *Unconventional Protein Secretion* 3-29 (Springer, 2016).
- 48 Araujo, T. L. *et al.* Protein disulfide isomerase externalization in endothelial cells 48 follows classical and unconventional routes. *Free Radical Biology and Medicine* **103**, 49 199-208 (2017).
- 49 Giordano, F. *et al.* PI (4, 5) P2-dependent and Ca2+-regulated ER-PM interactions 826 mediated by the extended synaptotagmins. *Cell* **153**, 1494-1509 (2013).
- 50 Zwicker, J. I. *et al.* Targeting protein disulfide isomerase with the flavonoid isoquercetin to improve hypercoagulability in advanced cancer. *JCI insight* **4** (2019).
- Jayachandran, M. *et al.* Isoquercetin upregulates antioxidant genes, suppresses
 inflammatory cytokines and regulates AMPK pathway in streptozotocin-induced
 diabetic rats. *Chemico-biological interactions* 303, 62-69 (2019).
- Stopa, J. D. *et al.* Protein disulfide isomerase inhibition blocks thrombin generation in
 humans by interfering with platelet factor V activation. *JCI insight* 2 (2017).
- Hynes, R. O. Integrins: bidirectional, allosteric signaling machines. *cell* 110, 673-687 (2002).
- 83654Cho, J. *et al.* Protein disulfide isomerase capture during thrombus formation in vivo837depends on the presence of β3 integrins. *Blood, The Journal of the American Society*838of Hematology 120, 647-655 (2012).
- 83955Swiatkowska, M., Szymański, J., Padula, G. & Cierniewski, C. S. Interaction and840functional association of protein disulfide isomerase with $\alpha V\beta 3$ integrin on841endothelial cells. *The FEBS journal* **275**, 1813-1823 (2008).
- 842 56 Passam, F. *et al.* Mechano-redox control of integrin de-adhesion. *Elife* 7, e34843 (2018).
- S. *et al.* The platelet integrin αIIbβ3 has an endogenous thiol isomerase
 activity. *Journal of Biological Chemistry* 275, 36984-36990 (2000).
- Lahav, J. *et al.* Enzymatically catalyzed disulfide exchange is required for platelet
 adhesion to collagen via integrin α2β1. *Blood* 102, 2085-2092 (2003).
- ⁸⁴⁸ 59 Inoue, O., Suzuki-Inoue, K., Dean, W. L., Frampton, J. & Watson, S. P. Integrin α2β1

- mediates outside-in regulation of platelet spreading on collagen through activation of
 Src kinases and PLCγ2. *The Journal of cell biology* 160, 769-780 (2003).
- 85160Wan, S. W. *et al.* Endothelial cell surface expression of protein disulfide isomerase852activates β 1 and β 3 integrins and facilitates dengue virus infection. Journal of cellular853biochemistry 113, 1681-1691 (2012).
- 85461Hahm, E. *et al.* Extracellular protein disulfide isomerase regulates ligand-binding855activity of α Mβ2 integrin and neutrophil recruitment during vascular inflammation.856Blood, The Journal of the American Society of Hematology 121, 3789-3800 (2013).
- 62 Gaspar, R. S. *et al.* Myricetin, the main flavonoid in Syzygium cumini leaf, is a novel
 inhibitor of platelet thiol isomerases PDI and ERp5. *Frontiers in pharmacology* 10,
 1678 (2020).
- Mangin, P. H. *et al.* Immobilized fibrinogen activates human platelets through
 glycoprotein VI. *haematologica* 103, 898 (2018).
- Remijn, J. A. *et al.* Role of ADP receptor P2Y12 in platelet adhesion and thrombus
 formation in flowing blood. *Arteriosclerosis, thrombosis, and vascular biology* 22,
 686-691 (2002).
- 865 65 Bergmeier, W., Chauhan, A. K. & Wagner, D. D. Glycoprotein Ibα and von
 866 Willebrand factor in primary platelet adhesion and thrombus formation: lessons from
 867 mutant mice. *Thrombosis and haemostasis* 99, 264-270 (2008).
- 868 66 Blenner, M. A., Dong, X. & Springer, T. A. Structural basis of regulation of von
 869 Willebrand factor binding to glycoprotein Ib. *Journal of Biological Chemistry* 289,
 870 5565-5579 (2014).
- K. et al. von Willebrand factor is dimerized by protein disulfide isomerase. *Blood, The Journal of the American Society of Hematology* 127, 1183-1191 (2016).
- Burgess, J. K. *et al.* Physical proximity and functional association of glycoprotein 1bα
 and protein-disulfide isomerase on the platelet plasma membrane. *Journal of Biological Chemistry* 275, 9758-9766 (2000).
- 876 69 Stopa, J. D., Baker, K. M., Grover, S. P., Flaumenhaft, R. & Furie, B. Kinetic-based
 877 trapping by intervening sequence variants of the active sites of protein-disulfide
 878 isomerase identifies platelet protein substrates. *Journal of Biological Chemistry* 292,
 879 9063-9074 (2017).
- Ki, J. *et al.* Platelet protein disulfide isomerase promotes glycoprotein ibα-mediated
 platelet-neutrophil interactions under thromboinflammatory conditions. *Circulation*139, 1300-1319 (2019).
- Rapaport, S. I. & Rao, L. V. M. The tissue factor pathway: how it has become a
 "prima ballerina". *Thrombosis and haemostasis* 73, 007-017 (1995).
- Ahamed, J. *et al.* Disulfide isomerization switches tissue factor from coagulation to cell signaling. *Proceedings of the National Academy of Sciences* 103, 13932-13937 (2006).
- Popescu, N. I., Lupu, C. & Lupu, F. Extracellular protein disulfide isomerase
 regulates coagulation on endothelial cells through modulation of phosphatidylserine
 exposure. *Blood* 116, 993-1001 (2010).
- Ansari, S. A., Pendurthi, U. R. & Rao, L. V. M. Role of cell surface lipids and thioldisulphide exchange pathways in regulating the encryption and decryption of tissue
 factor. *Thrombosis and haemostasis* 119, 860-870 (2019).
- 894 75 Versteeg, H. H. & Ruf, W. Tissue factor coagulant function is enhanced by protein895 disulfide isomerase independent of oxidoreductase activity. *Journal of Biological*896 *Chemistry* 282, 25416-25424 (2007).
- Chen, F. *et al.* Protein disulfide isomerase enhances tissue factor-dependent thrombin
 generation. *Biochemical and biophysical research communications* 501, 172-177

(2018). 899 77 Falati, S., Gross, P., Merrill-Skoloff, G., Furie, B. C. & Furie, B. Real-time in vivo 900 imaging of platelets, tissue factor and fibrin during arterial thrombus formation in the 901 mouse. Nature medicine 8, 1175-1180 (2002). 902 Robinson, R. M. et al. Inhibitors of the protein disulfide isomerase family for the 903 78 treatment of multiple myeloma. Leukemia 33, 1011-1022 (2019). 904 79 Delaney, M. K. et al. Differential Roles of the NADPH-Oxidase 1 and 2 in Platelet 905 Activation and Thrombosis. Arterioscler Thromb Vasc Biol 36, 846-854, 906 doi:10.1161/ATVBAHA.116.307308 (2016). 907 908 80 Bayraktutan, U., Blayney, L. & Shah, A. M. Molecular characterization and localization of the NAD (P) H oxidase components gp91-phox and p22-phox in 909 endothelial cells. Arteriosclerosis, thrombosis, and vascular biology 20, 1903-1911 910 911 (2000).81 Griendling, K. K., Minieri, C. A., Ollerenshaw, J. D. & Alexander, R. W. Angiotensin 912 II stimulates NADH and NADPH oxidase activity in cultured vascular smooth muscle 913 cells. Circulation research 74, 1141-1148 (1994). 914 915 82 Seno, T. et al. Involvement of NADH/NADPH oxidase in human platelet ROS production. Thrombosis research 103, 399-409 (2001). 916 Rastogi, R., Geng, X., Li, F. & Ding, Y. NOX activation by subunit interaction and 83 917 918 underlying mechanisms in disease. Frontiers in cellular neuroscience 10, 301 (2017). Vara, D., Campanella, M. & Pula, G. The novel NOX inhibitor 2-acetylphenothiazine 919 84 impairs collagen-dependent thrombus formation in a GPVI-dependent manner. Br J 920 921 *Pharmacol* **168**, 212-224, doi:10.1111/j.1476-5381.2012.02130.x (2013). Lambeth, J. D., Kawahara, T. & Diebold, B. Regulation of Nox and Duox enzymatic 922 85 activity and expression. Free Radical Biology and Medicine 43, 319-331 (2007). 923 924 86 Schroder, K., Weissmann, N. & Brandes, R. P. Organizers and activators: Cytosolic Nox proteins impacting on vascular function. Free Radic Biol Med 109, 22-32, 925 doi:10.1016/j.freeradbiomed.2017.03.017 (2017). 926 927 87 Vara, D., Cifuentes-Pagano, E., Pagano, P. J. & Pula, G. A novel combinatorial technique for simultaneous quantification of oxygen radicals and aggregation reveals 928 the activation of unexpected redox patterns in platelets by different 929 physiopathological Haematologica 104, 1879-1891, 930 stimuli. doi:10.3324/haematol.2018.208819 (2019). 931 Gaspar, R. S., Ferreira, P. M., Mitchell, J. L., Pula, G. & Gibbins, J. M. Platelet-932 88 derived extracellular vesicles express NADPH oxidase-1 (Nox-1), generate 933 934 superoxide and modulate platelet function. Free Radical Biology and Medicine 165, 395-400 (2021). 935 89 de, A. P. A. M. et al. Protein disulfide isomerase redox-dependent association with 936 937 p47(phox): evidence for an organizer role in leukocyte NADPH oxidase activation. JLeukoc Biol 90, 799-810, doi:10.1189/jlb.0610324 (2011). 938 90 Fernandes, D. C., Manoel, A. H., Wosniak, J., Jr. & Laurindo, F. R. Protein disulfide 939 isomerase overexpression in vascular smooth muscle cells induces spontaneous 940 preemptive NADPH oxidase activation and Nox1 mRNA expression: effects of 941 nitrosothiol exposure. Biochem 942 Arch Biophys **484**, 197-204, 943 doi:10.1016/j.abb.2009.01.022 (2009). Pescatore, L. A. et al. Protein disulfide isomerase is required for platelet-derived 91 944 growth factor-induced vascular smooth muscle cell migration, Nox1 NADPH oxidase 945 946 expression, and RhoGTPase activation. Journal of Biological Chemistry 287, 29290-947 29300 (2012). Hordijk, P. L. Regulation of NADPH oxidases: the role of Rac proteins. Circulation 948 92

- research 98, 453-462 (2006).
 Sousa, H. R. et al. Novel antiplatelet role for a protein disulfide isomerase-targeted
 peptide: evidence of covalent binding to the C-terminal CGHC redox motif. J Thromb *Haemost* 15, 774-784, doi:10.1111/jth.13633 (2017).
 Gimenez, M. et al. Redox Activation of Nox1 (NADPH Oxidase 1) Involves an
- 953 94 Gimenez, M. *et al.* Redox Activation of Nox1 (NADPH Oxidase 1) Involves an
 954 Intermolecular Disulfide Bond Between Protein Disulfide Isomerase and p47(phox) in
 955 Vascular Smooth Muscle Cells. *Arterioscler Thromb Vasc Biol* **39**, 224-236,
 956 doi:10.1161/ATVBAHA.118.311038 (2019).
- 957 95 Zhou, J. *et al.* The C-terminal CGHC motif of protein disulfide isomerase supports
 958 thrombosis. *J Clin Invest* 125, 4391-4406, doi:10.1172/JCI80319 (2015).
- 96 Gaspar, R. S. *et al.* Protein Disulphide Isomerase and NADPH Oxidase 1 Cooperate
 960 to Control Platelet Function and Are Associated with Cardiometabolic Disease Risk
 961 Factors. *Antioxidants* 10, 497 (2021).
- 96297Schiattarella, G. G. et al. Rac1 modulates endothelial function and platelet963aggregation in diabetes mellitus. Journal of the American Heart Association 7,964e007322 (2018).
- 965 98 Wang, X. *et al.* p47phox deficiency impairs platelet function and protects mice
 966 against arterial and venous thrombosis. *Redox Biology*, 101569 (2020).
- 967 99 Leavesley, D. I. *et al.* Vitronectin—master controller or micromanager? *IUBMB life*968 65, 807-818 (2013).
- Preissner, K. T. & Seiffert, D. Role of vitronectin and its receptors in haemostasis and vascular remodeling. *Thrombosis research* 89, 1-21 (1998).
- 101 Reheman, A. *et al.* Vitronectin stabilizes thrombi and vessel occlusion but plays a dual role in platelet aggregation. *Journal of Thrombosis and Haemostasis* 3, 875-883
 973 (2005).
- 102 Essex, D. W., Miller, A., Swiatkowska, M. & Feinman, R. D. Protein Disulfide
 105 Isomerase Catalyzes the Formation of Disulfide-Linked Complexes of Vitronectin
 106 with Thrombin– Antithrombin. *Biochemistry* 38, 10398-10405 (1999).
- Moretti, A. I. *et al.* Conserved gene microsynteny unveils functional interaction
 between protein disulfide isomerase and rho guanine-dissociation inhibitor families. *Scientific reports* 7, 1-18 (2017).
- Aslan, J. E. & McCarty, O. J. Rho GTPases in platelet function. Journal of Thrombosis and Haemostasis 11, 35-46 (2013).
- Sobierajska, K. *et al.* Protein disulfide isomerase directly interacts with β-actin
 Cys374 and regulates cytoskeleton reorganization. *Journal of Biological Chemistry* **289**, 5758-5773 (2014).
- 985106Versteeg, H. H. & Ruf, W. Tissue factor coagulant function is enhanced by protein-
disulfide isomerase independent of oxidoreductase activity. J Biol Chem 282, 25416-
25424, doi:10.1074/jbc.M702410200 (2007).
- 107 Chen, K., Detwiler, T. C. & Essex, D. W. Characterization of protein disulphide
 isomerase released from activated platelets. *Br J Haematol* 90, 425-431 (1995).
- Radomski, M., Palmer, R. & Moncada, S. An L-arginine/nitric oxide pathway present
 in human platelets regulates aggregation. *Proceedings of the National Academy of Sciences* 87, 5193-5197 (1990).
- Russo, I. *et al.* The activity of constitutive nitric oxide synthase is increased by the pathway cAMP/cAMP-activated protein kinase in human platelets. New insights into the antiaggregating effects of cAMP-elevating agents. *Thromb Res* 114, 265-273, doi:10.1016/j.thromres.2004.06.036 (2004).
- 110 Kwon, H. W., Shin, J. H., Cho, H. J., Rhee, M. H. & Park, H. J. Total saponin from
 Korean Red Ginseng inhibits binding of adhesive proteins to glycoprotein IIb/IIIa via

- phosphorylation of VASP (Ser(157)) and dephosphorylation of PI3K and Akt. J *Ginseng Res* 40, 76-85, doi:10.1016/j.jgr.2015.05.004 (2016).
- 1001111Fuentes, E. & Palomo, I. Role of oxidative stress on platelet hyperreactivity during1002aging. Life Sci 148, 17-23, doi:10.1016/j.lfs.2016.02.026 (2016).
- 1003 112 Ramachandran, N., Root, P., Jiang, X. M., Hogg, P. J. & Mutus, B. Mechanism of 1004 transfer of NO from extracellular S-nitrosothiols into the cytosol by cell-surface 1005 protein disulfide isomerase. *Proc Natl Acad Sci U S A* 98, 9539-9544, 1006 doi:10.1073/pnas.171180998 (2001).
- 1007 113 Root, P., Sliskovic, I. & Mutus, B. Platelet cell-surface protein disulphide-isomerase
 1008 mediated S-nitrosoglutathione consumption. *Biochem J* 382, 575-580,
 1009 doi:10.1042/BJ20040759 (2004).
- 1010 114 Peixoto, A. S. *et al.* Peroxynitrite preferentially oxidizes the dithiol redox motifs of
 1011 protein-disulfide isomerase. *J Biol Chem* 293, 1450-1465,
 1012 doi:10.1074/jbc.M117.807016 (2018).
- 1013 115 Shah, C. M., Bell, S. E., Locke, I. C., Chowdrey, H. S. & Gordge, M. P. Interactions between cell surface protein disulphide isomerase and S-nitrosoglutathione during nitric oxide delivery. *Nitric Oxide* 16, 135-142, doi:10.1016/j.niox.2006.08.001 (2007).
- 1017 116 Bell, S. E., Shah, C. M. & Gordge, M. P. Protein disulfide-isomerase mediates 1018 delivery of nitric oxide redox derivatives into platelets. *Biochem J* 403, 283-288, 1019 doi:10.1042/BJ20061146 (2007).
- 1020 117 Bekendam, R. H. *et al.* Protein disulfide isomerase regulation by nitric oxide
 1021 maintains vascular quiescence and controls thrombus formation. *Journal of* 1022 *Thrombosis and Haemostasis* 16, 2322-2335 (2018).
- 1023 118 Matsuo, Y. *et al.* Identification of a novel thioredoxin-related transmembrane protein.
 1024 *Journal of Biological Chemistry* 276, 10032-10038 (2001).
- 1025119Hoffstrom, B. G. et al. Inhibitors of protein disulfide isomerase suppress apoptosis1026induced by misfolded proteins. Nature chemical biology 6, 900-906 (2010).
- 1027 120 Kaplan, A. *et al.* Small molecule-induced oxidation of protein disulfide isomerase is
 1028 neuroprotective. *Proceedings of the National Academy of Sciences* 112, E2245-E2252
 1029 (2015).
- 1030121Vatolin, S. *et al.* Novel protein disulfide isomerase inhibitor with anticancer activity1031in multiple myeloma. *Cancer research* **76**, 3340-3350 (2016).
- 1032 122 Bekendam, R. H. *et al.* A substrate-driven allosteric switch that enhances PDI catalytic activity. *Nature communications* **7**, 1-11 (2016).
- 1034 123 Shergalis, A. *et al.* Characterization of aminobenzylphenols as protein disulfide
 1035 isomerase inhibitors in glioblastoma cell lines. *Journal of medicinal chemistry* 63,
 10263-10286 (2020).
- 1037 124 Kyani, A. *et al.* Discovery and mechanistic elucidation of a class of protein disulfide
 1038 isomerase inhibitors for the treatment of glioblastoma. *ChemMedChem* 13, 164-177
 1039 (2018).
- 1040 125 Sousa, H. *et al.* Novel antiplatelet role for a protein disulfide isomerase-targeted 1041 peptide: evidence of covalent binding to the C-terminal CGHC redox motif. *Journal* 1042 *of Thrombosis and Haemostasis* 15, 774-784 (2017).
- 1043126Zhou, J. et al. The C-terminal CGHC motif of protein disulfide isomerase supports1044thrombosis. The Journal of clinical investigation 125, 4391-4406 (2015).
- 1045 127 Cole, K. S. *et al.* Characterization of an A-site selective protein disulfide isomerase
 1046 A1 inhibitor. *Biochemistry* 57, 2035-2043 (2018).
- 1047 128 Xu, S. *et al.* Inhibition of protein disulfide isomerase in glioblastoma causes marked 1048 downregulation of DNA repair and DNA damage response genes. *Theranostics* **9**,

1049 2282 (2019).

- 1050 129 Cole, K. S. *et al.* Characterization of an A-Site Selective Protein Disulfide Isomerase
 1051 A1 Inhibitor. *Biochemistry* 57, 2035-2043, doi:10.1021/acs.biochem.8b00178 (2018).
- 130 Watashi, K. *et al.* Cyclophilin B is a functional regulator of hepatitis C virus RNA
 polymerase. *Molecular cell* 19, 111-122 (2005).
- 1054 131 Kaplan, A. & Stockwell, B. R. Structural elucidation of a small molecule inhibitor of 1055 protein disulfide isomerase. *ACS medicinal chemistry letters* **6**, 966-971 (2015).
- 1056 132 Chlebowska-Tuz, J. *et al.* Inhibition of protein disulfide isomerase induces differentiation of acute myeloid leukemia cells. *haematologica* 103, 1843-1852 (2018).
- 1059 133 Greve, E. *et al.* Route exploration and synthesis of the reported pyridone-based PDI inhibitor STK076545. *Organic & Biomolecular Chemistry* 18, 6665-6681, doi:10.1039/D0OB01205J (2020).
- 134 Xu, S. *et al.* Discovery of an orally active small-molecule irreversible inhibitor of
 protein disulfide isomerase for ovarian cancer treatment. *Proceedings of the National Academy of Sciences* 109, 16348-16353 (2012).
- 1065135Khodier, C. et al. in Probe Reports from the NIH Molecular Libraries Program1066[Internet](National Center for Biotechnology Information (US), 2014).
- 1067 136 Xiong, B., Jha, V., Min, J.-K. & Cho, J. Protein disulfide isomerase in cardiovascular
 1068 disease. *Experimental & Molecular Medicine*, 1-10 (2020).
- 1069 137 Kung, P.-H. *et al.* HPW-RX40 prevents human platelet activation by attenuating cell
 1070 surface protein disulfide isomerases. *Redox Biology* 13, 266-277 (2017).
- 1071 138 Holbrook, L. M. *et al.* Zafirlukast is a broad-spectrum thiol isomerase inhibitor that
 1072 inhibits thrombosis without altering bleeding times. *British Journal of Pharmacology*1073 (2020).
- 1074 139 Karala, A. R. & Ruddock, L. W. Bacitracin is not a specific inhibitor of protein disulfide isomerase. *The FEBS journal* 277, 2454-2462 (2010).
- 1076140Jasuja, R. et al. Protein disulfide isomerase inhibitors constitute a new class of1077antithrombotic agents. The Journal of clinical investigation 122, 2104-2113 (2012).
- 1078141Lin, L. et al. Quercetin-3-rutinoside inhibits protein disulfide isomerase by binding to1079its b' x domain. Journal of Biological Chemistry 290, 23543-23552 (2015).
- 1080142Zou, J. et al. Discovery of a novel ERp57 inhibitor as antiplatelet agent from danshen1081(Salvia miltiorrhiza). Evidence-Based Complementary and Alternative Medicine 20181082(2018).
- 1083 143 Giamogante, F. *et al.* Punicalagin, an active pomegranate component, is a new inhibitor of PDIA3 reductase activity. *Biochimie* **147**, 122-129 (2018).
- 1085 144 Ren, L. *et al.* Molecular docking-assisted screening reveals tannic acid as a natural
 1086 protein disulphide isomerase inhibitor with antiplatelet and antithrombotic activities.
 1087 Journal of Cellular and Molecular Medicine (2020).
- 1088145Kao, C.-C. *et al.* Juglone prevents human platelet aggregation through inhibiting Akt1089and protein disulfide isomerase. *Phytomedicine*, 153449 (2020).
- Wright, B., Watson, K. A., McGuffin, L. J., Lovegrove, J. A. & Gibbins, J. M. GRID
 and docking analyses reveal a molecular basis for flavonoid inhibition of Src family
 kinase activity. *The Journal of nutritional biochemistry* 26, 1156-1165 (2015).
- 1093 147 Wright, B. *et al.* A structural basis for the inhibition of collagen-stimulated platelet
 1094 function by quercetin and structurally related flavonoids. *British journal of* 1095 *pharmacology* 159, 1312-1325 (2010).
- 1096148Guerrero, J. et al. Flavonoids inhibit platelet function through binding to the1097thromboxane A2 receptor. Journal of Thrombosis and Haemostasis 3, 369-3761098(2005).

- 149 Lee, A. C.-L., Harris, J. L., Khanna, K. K. & Hong, J.-H. A comprehensive review on current advances in peptide drug development and design. *International journal of molecular sciences* 20, 2383 (2019).
- 1102 150 de A. Paes, A. M. *et al.* Protein disulfide isomerase redox-dependent association with
 p47phox: evidence for an organizer role in leukocyte NADPH oxidase activation.
 1104 *Journal of leukocyte biology* 90, 799-810 (2011).
- 1105 151 Chamberlain, N. *et al.* Lung epithelial protein disulfide isomerase A3 (PDIA3) plays
 1106 an important role in influenza infection, inflammation, and airway mechanics. *Redox*1107 *biology* 22, 101129 (2019).
- 1108 152 ZHAO, Z.-z., CHEN, F.-w., ZHOU, J.-s. & Cyrus, T. H. C. Effect of Small-molecule
 1109 Inhibitor of Protein Disulfide Isomerase-PACMA31 in Platelet Activation and
 1110 Thrombosis. *Chinese Journal of Thrombosis and Hemostasis*, 2 (2016).
- 1111 153 Cui, G. *et al.* Novel anti-thrombotic agent for modulation of protein disulfide
 1112 isomerase family member ERp57 for prophylactic therapy. *Scientific Reports* 5, 10353 (2015).
- 1114 154 Trnková, L., Ricci, D., Grillo, C., Colotti, G. & Altieri, F. Green tea catechins can
 1115 bind and modify ERp57/PDIA3 activity. *Biochimica et Biophysica Acta (BBA)*1116 *General Subjects* 1830, 2671-2682 (2013).
- 1117 155 Mosawy, S., Gaiz, A., Karaksha, A. & Singh, I. The green tea extract epigallocatechin
 1118 gallate inhibits human platelet function but not plasma coagulation. *International*1119 *Journal of Prevention and Treatment* 5, 17-21 (2016).
- 1120 156 Khan, M. M. *et al.* Discovery of a small molecule PDI inhibitor that inhibits reduction
 1121 of HIV-1 envelope glycoprotein gp120. *ACS chemical biology* 6, 245-251 (2011).
- 1122 157 Sachetto, A. T. A., Rosa, J. G. & Santoro, M. L. Rutin (quercetin-3-rutinoside)
 1123 modulates the hemostatic disturbances and redox imbalance induced by Bothrops
 1124 jararaca snake venom in mice. *PLoS neglected tropical diseases* 12, e0006774 (2018).
- 1125 158 Klappa, P., Hawkins, H. C. & Freedman, R. B. Interactions between protein disulphide isomerase and peptides. *European journal of biochemistry* 248, 37-42 (1997).
- 1128 159 Ozaki, Y. *et al.* Mastoparan, a wasp venom, activates platelets via pertussis toxin-1129 sensitive GTP-binding proteins. *Biochemical and biophysical research* 1130 *communications* **170**, 779-785 (1990).
- 1131