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Drugs Used in Thromboembolic Disorders: An Insight Into Their Mechanisms

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Abstract:

9 In the United States alone, more than 6 million patients receive long-term anti-platelet and
10 anticoagulation therapy. The hemostatic system must maintain a balance between fibrin
11 formation (coagulation) and fibrin dissolution (fibrinolysis). Thrombin and Factor Xa are
12 two of the most important components of the coagulation cascade. Any disruption in this
13 cascade can lead to either thrombosis, hemorrhage or both. The clotting process is a dynamic,
14 highly interwoven array of multiple processes. There are four different phases involved in the
15 response of activated platelets: Adhesion, aggregation, secretion and the procoagulant
16 activity. The coagulation cascade is a coordinated sequence of linked enzymatic reactions in
17 which each reaction product converts the subsequent inactive zymogen into an active serine
18 protease that are responsible for the conversion of soluble plasma fibrinogen into insoluble
19 fibrin. Several antithrombotic factors regulate coagulation and limit the production of
20 thrombin to prevent the perpetuation of coagulation and thrombus formation; these include
21 protein C/protein S, antithrombin, heparin cofactor and tissue factor pathway inhibitor.
22 Antiplatelet agents play a major role in the management of cerebrovascular, peripheral
23 vascular and cardiovascular diseases. Aspirin is a non-steroidal anti-inflammatory drug that
24 works by irreversibly inhibiting cyclo-oxygenase 1 and 2 by covalent acetylation. Cilostazol,
25 a specific and strong inhibitor of PDE3 in platelets and smooth muscle cells was approved in
26 the USA in 1999 for the treatment of intermittent claudication. Dipyridamole affects platelet
27 function by inhibiting the reuptake of adenosine by red blood cells, in this way enhancing
28 plasma levels of this vasodilator and platelet inhibitory nucleoside; it acts as an inhibitor of
29 PDE5 and PDE3, thus increasing intraplatelet cAMP and/or cGMP; and it also acts as an
30 antioxidant by scavenging free radicals that inactivate cyclo-oxygenase, thus enhancing PGI₂
31 biosynthesis. Vorapaxar, a first in its class, is an orally available PAR-1 antagonist approved
32 for patients with prior myocardial infarction or peripheral arterial disease with no previous
33 history of stroke or TIA, and is added to standard therapy for long-term secondary
34 prevention of thrombotic CV events. Abciximab, eptifibatide, tirofiban all bind the
35 glycoprotein receptor IIb/IIIa on activated platelets, preventing aggregation. Warfarin is a
36 Vitamin K antagonist that interferes with γ -carboxylation of vitamin K-dependent clotting
37 factors II, VII, IX, and X, and proteins C and S. Dabigatran is a potent, competitive inhibitor
38 of thrombin, while Apixaban, edoxaban and rivaroxaban all selectively inhibit factor Xa.
39 Heparins, acts indirectly by binding to anti-thrombin (AT) rather than acting directly on the
40 coagulation factors. This interaction converts AT to a rapid inactivator of factor IIa and factor
41 Xa. Unfractionated heparin, LMW heparin (enoxaparin, dalteparin) and fondaparinux all

42 inactivate factor Xa, but unfractionated heparin is a much more efficient inactivator of
43 thrombin. Drugs such as anti-platelet therapy and anti-coagulants are frequently used in
44 clinical settings. It is imperative that the physicians have a thorough understanding of these
45 agents. An insight into the mechanism of how these medications act serves as a prelude to
46 understanding to the pharmacology of these drugs.

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48

49 **Introduction:**

50

51 Cardiovascular diseases like myocardial infarction, stroke, deep-venous
52 thrombosis, and pulmonary embolism are among the leading causes of mortality
53 worldwide [1]. In the United States alone, more than 6 million patients receive
54 long-term anti-platelet and anticoagulation therapy for the prevention of various
55 causes of thromboembolism that include venous thromboembolism, atrial
56 fibrillation or placement of a mechanical heart-valve prosthesis [2].

57 Venous thromboembolism (VTE) comprising deep venous thrombosis
58 (DVT) and pulmonary embolism (PE) is the third vascular cause of death after
59 myocardial infarction and stroke and is a significant cause of morbidity and
60 mortality. Annual incidence of VTE is estimated to be approximately 1 or 2
61 cases per 1000 persons in the general population [3]. however, VTE is often
62 misdiagnosed, asymptomatic and unrecognized at death and regular post-
63 mortem examinations are missing. These factors are believed to result in
64 marked underestimations of VTE incidence. Complications that arise after VTE
65 including chronic thromboembolic pulmonary hypertension and post-thrombotic
66 syndrome account for the substantial increase in morbidity and healthcare costs
67 incurred from VTE.

68 In order to ensure adequate perfusion of blood through tissues and to
69 prevent the free flow of blood at sites of injury, the hemostatic system must
70 maintain a balance between fibrin formation (coagulation) and fibrin dissolution
71 (fibrinolysis) [4]. The major structural component of blood clot is fibrin, which
72 is formed by the proteolytic action of thrombin on fibrinogen. Fibrinolysis is a
73 process by which fibrin is degraded into fibrin degradation products by the
74 action of plasmin. Thrombin and Factor Xa are two of the most important
75 components of the coagulation cascade [5]. Any disruption in this cascade can
76 lead to either thrombosis, hemorrhage or both.

77 Anticoagulants include a variety of agents that work by inhibiting one or
78 more steps in the coagulation cascade. Their mechanisms include direct
79 enzymatic inhibition, indirect inhibition by binding to antithrombin, antagonism
80 of vitamin K-dependent coagulation factors by either the modification of their
81 calcium-binding properties or by preventing their synthesis in the liver.

82 The steadily expanding list of antiplatelet and anticoagulants provide a
83 wide range of agents for prevention and management of thromboembolic
84 diseases. However, the appropriate use of these agents require an understanding
85 of their individual characteristics, benefits and risks. This article provides a
86 review of the recent advances in the use of antiplatelet and anticoagulant
87 therapies and an insight into their mechanisms in the coagulation cascade.
88

89 **Mechanisms of action**

90 *Normal haemostasis*

91 The hemostatic system provides a balance between procoagulant and
92 anticoagulant forces. When a blood vessel wall is disrupted, the hemostatic
93 response must be quick, localized and carefully regulated. Any disturbances in
94 this pathway may lead to either abnormal bleeding or thrombosis.

95 The clotting process is a dynamic, highly interwoven array of multiple
96 processes [6]. The phases of the process include:

- 97 1. Endothelial injury and formation of the platelet plug
- 98 2. Propagation of the clotting process by the coagulation cascade
- 99 3. Termination of clotting by antithrombotic control mechanisms.
- 100 4. Removal of the clot by fibrinolysis.

102 *Formation of a platelet plug*

103 In order to stop bleeding, the initial hemostatic response is the formation of a
104 platelet plug at the site of vascular injury. Endothelial cell activation and the
105 exposure of the subendothelial elements may promote recruitment of platelets,
106 procoagulant factors and other cell types [7].

107 The activation of platelets is done through a number of physiological stimuli
108 that include adenosine diphosphate (ADP), thrombin, collagen and epinephrine.
109 The summary of all the important steps involved in formation of a platelet plug
110 has been mentioned below in the Figure 1.

111 The adherence of platelets is prevented by the secretion of prostacyclin and
112 nitric oxide from the intact endothelium. Endothelial injury impairs this process
113 and leads to the exposure of subendothelial elements such as collagen that
114 activates platelets. The two most important platelet collagen receptors are
115 GPIa/IIa (integrin- $\alpha 2\beta 1$) and GPVI [8].

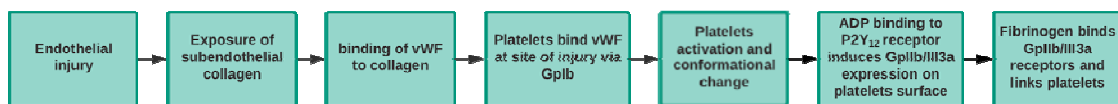
116 The principle receptors for thrombin on human platelets are a family of G-
117 protein coupled protease-activated receptors (PAR). PAR-1 is a high-affinity
118 receptor that mediates the effect of thrombin at low concentrations, while PAR-
119 4 is a low-affinity receptor that requires high levels of thrombin for activation.
120 Notably the binding of thrombin to PAR-1, which represents a very potent

121 platelet activation pathway, is necessary for thrombus formation but may not be
122 required for hemostasis [9, 10]. Vorapaxar is an oral PAR-1 antagonist
123 developed as an antiplatelet agent [11].

124 ADP released from damaged red blood cells and vessels activates the integrin
125 GPIIb-IIIa and subsequent binding of fibrinogen that induces platelet
126 aggregation. ADP mediates its action through binding of two purinergic G-
127 protein coupled receptors, P2Y1 and P2Y12 [12]. The P2Y1 receptor couples to
128 Gq and mobilizes intracellular calcium and mediate platelet shape change and
129 rapidly reversible aggregation; P2Y12 is coupled to a Gi that mediates a more
130 stable platelet aggregation [13]. The thienopyridine derivatives, clopidogrel and
131 ticlopidine are antiplatelet agents that inhibit the platelet aggregation induced by
132 ADP, thereby reducing ischemic events [14].

133 There are four different phases involved in the response of activated platelets:
134 Adhesion, aggregation, secretion and the procoagulant activity. The first
135 response of activated platelets is the deposition of platelets on the
136 subendothelial matrix followed by aggregation which is mediated via platelet-
137 platelet cohesion and the secretion of platelet granule proteins and finally the
138 enhancement of thrombin generation [15].

139
140



141

142 **Figure 1.** Summary of all the important steps in the formation of the platelet
143 plug.

144

145 *The coagulation Cascade*

146 The coagulation cascade is a coordinated sequence of linked enzymatic
147 reactions in which each reaction product converts the subsequent inactive
148 zymogen into an active serine protease that are responsible for the conversion of
149 soluble plasma fibrinogen into insoluble fibrin [16].

150 The primary initiating factor for the coagulation cascade is the exposure of
151 tissue factor at areas of endothelial or tissue injury to blood components and the
152 interaction of activated factor VII (FVIIa) with exposed tissue factor (TF).
153 Tissue factor may be found in leukocytes or platelets or in bloodborne cell-
154 derived microparticles. The TF-FVIIa complex then activates factors IX (FIX)
155 and X (FX), leading to the release and activation of factor V (FV) from platelet
156 α -granules. Activated FX (FXa) links with activated FV (FVa) converting factor

157 II (FII, also known as prothrombin) to thrombin (FIIa). Thrombin then converts
158 fibrinogen to fibrin and overlapping fibrin strands are cross-linked and
159 stabilized by activated factor XIII (FXIIIa) [17].

160 Several antithrombotic factors regulate coagulation and limit the production of
161 thrombin to prevent the perpetuation of coagulation and thrombus formation;
162 these include protein C/protein S, antithrombin, heparin cofactor and tissue
163 factor pathway inhibitor (TFPI) [18].

164 *The cell-based model of fibrin formation*

165
166 The cell-based model of hemostasis replaces the traditional hypothesis of
167 "cascade" and suggests that coagulation takes place in four overlapping steps on
168 distinct cell surfaces: initiation, amplification and propagation [19].

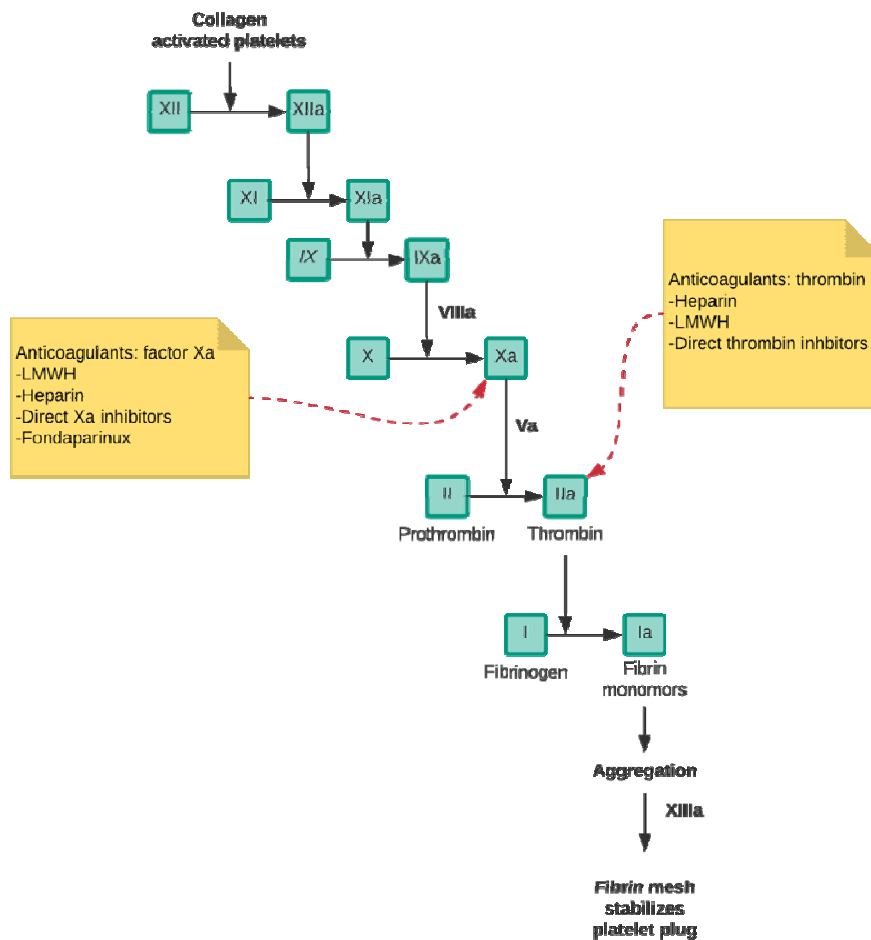
169
170 The initiation phase occurs when cells on the site of injury that express
171 TF on their surface are exposed to blood components. It results in a small
172 amount of FIXa and thrombin being generated that diffuses to the platelet from
173 the TF-bearing cell surface.

174 In The amplification phase, binding of thrombin to platelet surface
175 receptors results in extreme modifications in the platelet surface, leading to
176 changes in shape, release of granules and shuffling of membrane phospholipids
177 to generate procoagulant membrane surface. These modifications result in
178 activations of platelets that release vWF and generate FV, FVIII and FXI.

179 The propagation phase is characterized by the migration of large numbers
180 of platelets to the injury site. The different enzymes generated in earlier phases
181 assemble to form the tenase complex and the prothrombinase resulting in FXa
182 and thrombin generation (20).

183

184 Different anticoagulant agents work on this pathway; these include Heparin,
185 LMWH, direct thrombin inhibitors, direct Xa inhibitors, fondaparinux and
186 thrombolytics such as alteplase. the sites of action of the different
187 anticoagulants is described in the Figure 2.



188

189 **Figure 2.** Illustrates and summarizes the sites of action of different therapeutic
 190 agents.

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195 **Antiplatelet Agents**

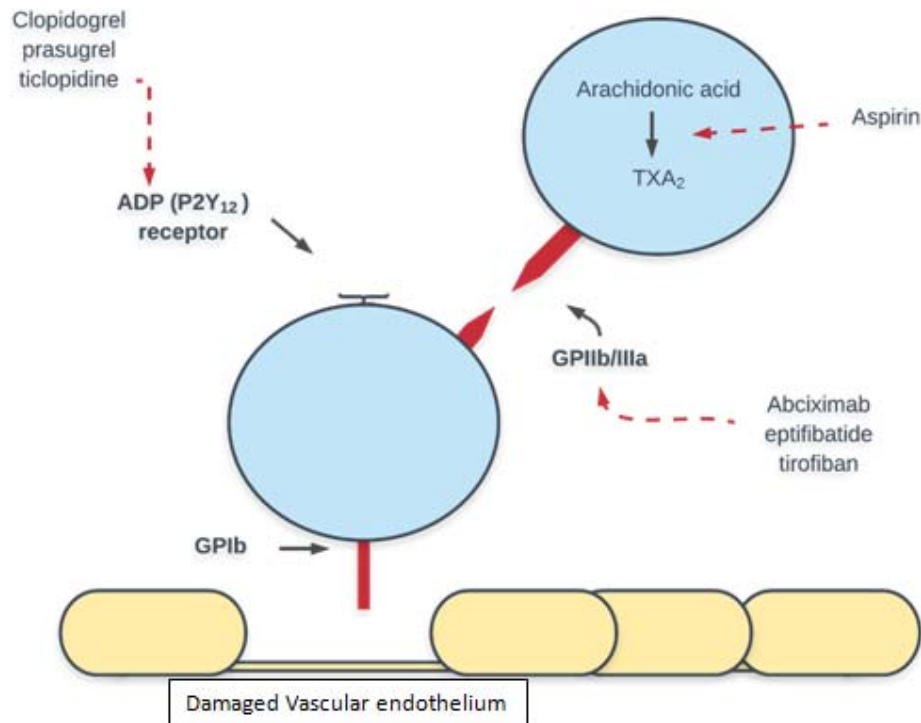
196 Antiplatelet agents play a major role in the management of cerebrovascular,
 197 peripheral vascular and cardiovascular diseases. Depending on the specific
 198 clinical indication and the patient's risk for thromboembolic or bleeding events,
 199 these agents can be used as monotherapy or combination therapy. A list of the
 200 currently available anti-platelet agents is provided in the Table 1.

201

202 **Table 1.** List of the available anti-platelet medications (generic and brand
203 names).
204

Drug generic name	Drug brand name
Aspirin / extended-release dipyridamole	Aggrenox
Aspirin / omeprazole	Yosprala
Aspirin extended-release	Durlaza
Cilostazol	Pletaal
Clopidogrel	Plavix
Dipyridamole	Persantine
Ticagrelor	Brilinta
Prasugrel	Effient
Ticlopidine	Ticlid
Vorapaxar	Zontivity
Abciximab	Reopro
Eptifibatide	Integrilin

205
206



207

208 **Figure 3.** Illustration of mechanism of action of different anti-platelet agents

209

210 **Aspirin**

211 Aspirin is a non-steroidal anti-inflammatory drug that works by irreversibly
 212 inhibiting cyclo-oxygenase 1 and 2 by covalent acetylation. This leads to
 213 decreased synthesis of prostaglandins and thromboxane A₂ particularly in the
 214 platelets. Thromboxane A₂ is an important mediator that is essential for platelet
 215 aggregation and therefore aspirin acts as an anti-aggregator of the platelets [18].
 216 Mechanism of action of various Antiplatelet agents has been described in the
 217 Figure 3.

218 The therapeutic effects of aspirin are dose dependent. Antithrombotic effects of
 219 aspirin have been reported with low doses (<300 mg/day), antipyretic, analgesic
 220 effects are seen with Intermediate doses (300–2400 mg/day) and anti-
 221 inflammatory effects are seen at High doses (2400–4000 mg/day) [21].

222 Potential side effects when aspirin is used as an anti-platelet medication include
 223 gastric erosion, gastric and duodenal ulceration. Side effects that are rarely seen
 224 when used as anti-platelet medication include nephropathy due to reduced blood
 225 flow to the kidneys, interstitial nephritis, hepatotoxicity, asthma, and GI
 226 bleeding [22].

227

228

229 *Cilostazol and dipyridamole*

230 Inhibition of platelet aggregation can be achieved by inhibiting certain
231 intracellular signalling pathways. Cyclic adenosine monophosphate (cAMP) and
232 cyclic guanosine monophosphate (cGMP) are two critical intracellular second
233 messengers with strong inhibitory activity on fundamental platelet functions.
234 These second messengers (cAMP and cGMP) are both metabolized by the class
235 of enzymes known as Phosphodiesterases (PDEs), which catalyzes the
236 hydrolysis of cAMP and cGMP, thus regulating platelet function.

237 Platelets possess three PDE isoforms (PDE2, PDE3 and PDE5), with different
238 selectivity for cAMP and cGMP. While PDE2 catalyzes the hydrolysis of both
239 cAMP and cGMP (cAMP = cGMP), PDE3 preferentially catalyzes the
240 hydrolysis of cAMP over cGMP (cAMP > cGMP) and PDE5 catalyzes the
241 hydrolysis of cGMP. The inhibition of PDEs may therefore exert a strong
242 platelet inhibitory effect.

243 Cilostazol, a specific and strong inhibitor of PDE3 in platelets and smooth
244 muscle cells was approved in the USA in 1999 for the treatment of intermittent
245 claudication. The overall effect of the drug is that it diminishes intracellular
246 calcium, causing smooth muscle cell relaxation and inhibition of platelet
247 activation. Cilostazol inhibits adenosine uptake, thus enhancing adenosine
248 levels that in turn enhance intracellular cAMP, resulting in additional increases
249 in cAMP. Cilostazol enhances the antiplatelet effects of the endothelium-
250 derived prostacyclin (PGI₂), which inhibits thrombosis in vivo and inhibits
251 shear, stress-induced platelet aggregation in vitro and ex vivo. One potential
252 benefit of the use of cilostazol over conventional antiplatelet therapy is the
253 relatively short recovery time of platelet function. The most common adverse
254 effects of cilostazol are headache, tachycardia, palpitations, soft stools and
255 diarrhoea [23-25].

256 Dipyridamole affects platelet function by acting on the following different
257 targets: it inhibits the reuptake of adenosine by red blood cells, in this way
258 enhancing plasma levels of this vasodilator and platelet inhibitory nucleoside; it
259 acts as an inhibitor of PDE5 and PDE3, thus increasing intraplatelet cAMP
260 and/or cGMP; and it acts as an antioxidant by scavenging free radicals that
261 inactivate cyclo-oxygenase, thus enhancing PGI₂ biosynthesis. Combination
262 therapy of aspirin and dipyridamole is more effective than aspirin alone in the
263 prevention of new serious vascular events in patients after non-disabling
264 cerebral ischaemia of presumed arterial origin [24, 26].

265 *ADP receptor inhibitors*

266 Red blood cells, damaged endothelial cells and activated platelets release ADP,
267 which is a potent activator of platelets which induces its adhesion and
268 aggregation. The response of the platelets to ADP is mediated by a family of

269 membrane bound receptors called P2 receptors. ADP binds to the P2Y₁ and
270 P2X₁ receptors leading to platelet aggregation. The binding of ADP to its Gi -
271 coupled P2Y₁₂ receptor liberates the Gi protein subunits G α and G $\beta\gamma$ and this
272 results in stabilisation of platelet aggregation. The subunit G α leads to inhibition
273 of adenylyl cyclase (AC), which reduces cyclic adenosine monophosphate
274 (cAMP) levels. This is also known to cause glycoprotein IIb/IIIa receptor
275 activation resulting in aggregation of platelets. Furthermore, G $\beta\gamma$ is known to
276 independently cause platelet aggregation via phosphatidylinositol 3-kinase
277 pathway, which can cause platelet aggregation.

278 The thienopyridine compounds ticlopidine and clopidogrel were the first ADP
279 inhibitors to be used clinically as antithrombotic drugs. Other agents in this
280 category include; prasugrel and ticagrelor. These agents inhibit platelet
281 aggregation by blocking the P2Y₁₂ receptor thus preventing the expression of
282 glycoprotein IIb/IIIa on platelets surface. Ticlopidine and clopidogrel are
283 effective antiplatelet agents and are useful in the prevention of stroke,
284 myocardial infarction, and vascular death in patients with vascular disease.
285 However, the major concern regarding safety of ticlopidine is severe and
286 sometimes fatal blood dyscrasias which is not observed with Clopidogrel [27,
287 28].

288

289 *Vorapaxar*

290 Thrombin is a potent platelet activator which acts via cell-surface protease-
291 activated receptors (PARs) which are widely expressed in human platelets.
292 Platelet activation through PAR-1 signalling results in extracellular ADP
293 release, resulting in activation of platelet ADP receptors, thereby promoting
294 aggregation of platelets. The PAR-1 receptor blockade produces potent
295 antiplatelet effects without affecting the ability of thrombin to generate fibrin
296 and without inhibiting platelet activation by collagen.

297 Vorapaxar, a first in its class, is an orally available PAR-1 antagonist approved
298 for patients with prior myocardial infarction or peripheral arterial disease and is
299 added to standard therapy for long-term secondary prevention of thrombotic CV
300 events. However, the use of this medication is associated with an increased risk
301 for bleeding due to which the prescribing information of the drug carries a
302 boxed warning for the same. Further, it is contraindicated in patients who have
303 history of stroke, TIA, or intracranial haemorrhage, because of an increased risk
304 for intracranial haemorrhage in these patient populations. [29].

305

306 *Glycoprotein IIb/IIIa inhibitors*

307 Abciximab, eptifibatide, tirofiban all bind the glycoprotein receptor IIb/IIIa on
308 activated platelets, preventing aggregation. Glycoprotein IIb/IIIa inhibitors have

309 been shown to reduce cardiac complications in patients undergoing
310 percutaneous coronary intervention and also to reduce the mortality in patients
311 with acute coronary syndromes who are not routinely scheduled for early
312 revascularization [30].

313 **Anticoagulant Agents:**

314 There are a variety of parenteral and oral anticoagulants available for the
315 management of thromboembolic disease. Parenteral agents such as low
316 molecular weight heparin (LMWH), unfractionated heparin (UFH) and
317 fondaparinux are used for initial, and sometimes longer term, treatment. On the
318 other hand Vitamin K antagonists are used for long-term treatment. The
319 available oral antagonists consist of warfarin, rivaroxaban, dabigatran,
320 edoxaban and apixaban. A list of available anticoagulants have been mentioned
321 in Table 2 [31, 32].

322

323 **Table 2:** List of available anti-coagulants, their generic name, brand name and
324 route of administration (ROA).

325

326

Drug generic name	Drug brand name	ROA
Heparin		Intravenous/Subcutaneous
Fondaparinux	Arixtra	Intravenous/Subcutaneous
Dalteparin	Fragmin	Intravenous/Subcutaneous
Enoxaparin	Lovenox	Intravenous/Subcutaneous
Warfarin	Coumadin	Oral
Rivaroxaban	Xarelto	Oral
Dabigatran	Pradaxa	Oral
Edoxaban	Savaysa	Oral
Apixaban	Eliquis	Oral

327

328 **Vitamin K antagonists (VKA)**

329 Warfarin is the most widely used anticoagulant in the world. It is most
330 commonly used to prevent embolic strokes in patients with atrial fibrillation.

331 Warfarin is a Vitamin K antagonist that interferes with γ -carboxylation of
332 vitamin K–dependent clotting factors II, VII, IX, and X, and proteins C and S.

333 The main adverse effect associated with warfarin is bleeding. The patients on
334 warfarin therapy are monitored regularly by measuring International
335 Normalized Ratio (INR). The target range for INR in patients is between 2 and
336 3. An INR over 3 increases the risk for bleeding whereas an INR below the
337 level of 2 increases the risk for thrombosis. Warfarin's narrow therapeutic index
338 makes it difficult to maintain this predefined target range of INR. This problem
339 is additionally compounded by the individual and pharmacogenomic variations
340 that effect the warfarin metabolism as well as the drug interactions associated
341 with the therapy. Polymorphisms in the genes encoding CYP2C9, (Hepatic
342 cytochrome that metabolizes warfarin) and VKORC1, (vitamin K epoxide
343 reductase, an enzyme that warfarin inhibits) have been shown to be the major
344 determinants of warfarin dosage. Other determinants of warfarin dosage include
345 dietary vitamin K intake, herbal products and dietary supplements which can
346 have a significant effect on the degree of anticoagulation.

347 Warfarin also interferes with the formation of γ -carboxyglutamate proteins that
348 is an essential part of osteocalcin, a bone regulating protein. This effect
349 contributes to the teratogenic effects ranging from skeletal to cartilaginous
350 malformations [33, 34, 35].

351

352 **Direct oral anticoagulants**

353 The treatment of venous thromboembolism (VTE) traditionally consists of
354 initial low-molecular-weight heparin (LMWH) or unfractionated heparin
355 followed by warfarin for at least 3 months. Recently, the development and
356 introduction of direct oral anticoagulants have proven to be an effective and safe
357 alternative [36]. These agents work by inhibiting factor Xa and IIa (thrombin) in
358 the coagulation cascade leading to inhibition of fibrin formation. Dabigatran is a
359 potent, nonpeptidic small molecule that specifically and reversibly inhibits both
360 free and clot-bound thrombin by binding to the active site of the thrombin
361 molecule, while Apixaban, edoxaban and rivaroxaban all selectively inhibit
362 factor Xa.

363 Advantages of direct oral anticoagulants, unlike Vitamin K antagonists (VKA)
364 like warfarin is that their therapy does not require frequent laboratory
365 monitoring and subsequent dose adjustments, which represents a major shift
366 from the traditional VKA-based therapies. Direct oral anticoagulants for the
367 treatment and prevention of thromboembolic disorders represent a major shift
368 from the traditional VKA-based therapies. Four direct oral anticoagulants
369 apixaban, dabigatran, edoxaban and rivaroxaban have completed large phase III
370 clinical studies and have been currently approved by the FDA for treatment and
371 prophylaxis of various thromboembolic conditions. These direct oral
372 anticoagulants bind directly to key proteins of the clotting cascade, leading to
373 the inhibition of fibrin formation.

374

375 The Thrombin clotting time assay evaluates thrombin activity in a plasma
376 sample directly and thus gives a direct measure of DTI activity. TT tests in
377 many hospital laboratories are readily available. The TT is particularly
378 sensitive to dabigatran effects and shows a linear dose response over therapeutic
379 concentrations [37].

380 However, with direct oral anticoagulants, severe bleeding may happen.
381 Furthermore, patients taking NOACs may experience trauma and may need
382 urgent surgery or procedures. Therefore, the accessibility of particular reversal
383 agents for NOACs in these emergency scenarios could enhance patient
384 management. Idarucizumab is a monoclonal humanized antibody fragment
385 created as a particular dabigatran reversal agent. Studies in vitro and ex vivo
386 have shown that idarucizumab quickly restores parameters of dabigatran-
387 prolonged coagulation to baseline values [38].

388

389

390

391 **Heparin and LMW heparin**

392 Heparins, including unfractionated heparin and a variety of low molecular
393 weight (LMW) heparin products have been used for over eighty years as an
394 anticoagulant in treatment and prophylaxis of venous thromboembolism,
395 Coronary Artery Disease, Unstable Angina and Non-Q-Wave Myocardial
396 infarction, Acute Myocardial infarction, Coronary Thrombolysis, Coronary
397 Angioplasty, Atrial Fibrillation, etc. These agents act indirectly by binding to
398 anti-thrombin (AT) rather than acting directly on the coagulation factors.

399 This interaction converts AT to a rapid inactivator of factor IIa and factor Xa
400 [39]. The antithrombin-mediated anticoagulant effects of heparin are best
401 described for factors Xa and thrombin, but a third factor, IXa, is an important
402 contributor. Factor XIa, a less important contributor to the anticoagulant action
403 of heparin / antithrombin, is structurally distinct from other proteases of
404 coagulation, but like factors IXa, Xa and thrombin it is created by a pro-enzyme
405 cleavage and is subject to heparin-bound antithrombin inhibition. Although
406 thrombin inhibition is a desirable therapeutic effect of unfractionated heparin,
407 nonspecific interactions with other proteins, such as platelet factor 4, result in
408 side effects that can be life-threatening [40].

409

410 Unfractionated heparin, LMW heparin (enoxaparin, dalteparin) and
411 fondaparinux all inactivate factor Xa, but unfractionated heparin is a much more
412 efficient inactivator of thrombin [41, 42]. Unfractionated heparin generally is
413 administered intravenously as an initial bolus followed by a continuous infusion

414 using a heparin nomogram while LMW heparin is administered subcutaneously
415 in fixed or weight-based dosing without monitoring [42].

416 The limitations of heparin use include osteopenia and heparin-induced
417 thrombocytopenia (HIT).

418 HIT is a life-threatening disorder that results from exposure to non-fractionated
419 or (less frequently) low-molecular heparin. Classically patients present with low
420 platelet counts (< 150,000 per cubic millimeter) or a relative decrease from
421 baseline by 50 percent or more. HIT is caused by antibodies against complexes
422 of platelet factor 4 (PF4) and heparin. The risk of thrombosis in patients with
423 HIT is more than 30 times that in control populations. The risk of thrombosis
424 remains high for days to weeks after heparin has been discontinued, even after
425 platelet counts have normalized. HIT occurs more frequently in patients treated
426 with low-molecular-weight heparin who have recently had exposure to
427 unfractionated heparin (within 100 days) than in those who have not had a
428 recent exposure to unfractionated heparin [44].

429

430 Osteopenia is caused as a result of binding of heparin to osteoblasts, which then
431 release factors that activate osteoclasts. The dosage regimen of the heparins are
432 calculated based on the body weight and therefore there is a possibility of an
433 inadvertent overexposure. A number of techniques have been published to
434 standardize the management of IV heparin treatment, including nomograms for
435 heparin dose adjustment and computer algorithms. A weight-based nomogram
436 using a starting dose of 18 U/kg/h heparin infusion (1,260 U/h for a 70-kg
437 patient) reduced recurrent thromboembolism in a randomized controlled trial
438 [45].

439 In case of Unfractionated heparin, Protamine sulfate is used as an antidote to
440 counter the excess heparin. However there are no clear guidelines for
441 management of toxicity of LMW heparins [46, 47]

442

443 **Conclusion:**

444 Cardiovascular events such as myocardial infarction, stroke, deep-venous
445 thrombosis, and pulmonary embolism are the most common life-threatening
446 conditions that are required to be treated on a timely basis and therefore prompt
447 recognition and management is vital. Drugs such as anti-platelet therapy and
448 anti coagulants are frequently used in clinical settings. It is imperative that the
449 physicians have a thorough understanding of these agents. An insight into the
450 mechanism of how these medications act serves as a prelude to understanding to
451 the pharmacology of these drugs.

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