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**Management of antiplatelet or** **anticoagulant therapy in endoscopy: A review of literature**

Maida M *et al.* Management of anticoagulants in endoscopy

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 **Abstract**

Endoscopic procedures hold a basal risk of bleeding that depends on the type of the procedure and patients’ comorbidities. Moreover, they are more often performed in patients taking antiplatelet and anticoagulants agents, increasing the potential risk of intraprocedural and delayed bleeding. On the other hand, the discontinuation of therapy may increase the cardiovascular risk when therapy is suspended or replaced. Therefore, it is of fundamental for each endoscopist to be aware of the bleeding risk for every procedure, in order to measure the risk-benefit ratio for each individual patient. Moreover, it is essential to know how to handle antiplatelet and anticoagulant agents before the procedure, as well the adequate timing for the resumption of therapy after endoscopy. The aim of this review is to analyze current evidence from literature assessing, for each procedure, the basal risk of bleeding and the risk of bleeding in patients taking anticoagulant, as well as to review the recommendation of American society for gastrointestinal endoscopy, European society of gastrointestinal endoscopy and British society of gastroenterology and Asian pacific association of gastroenterology and Asian pacific society for digestive endoscopy guidelines for the management of anticoagulants in urgent and elective endoscopic procedures.

**Key words:** Antiplatelet; Anticoagulant; Endoscopy; Management; Bleeding; Risk

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**Core tip:** Endoscopic procedures hold a basal risk of bleeding and they are more performed in patients taking antiplatelet and anticoagulant agents, increasing the potential risk of intraprocedural and delayed bleeding. The aim of this review is to analyze current evidence from literature assessing, for each procedure, the basal bleeding risk and the risk of bleeding in patients taking anticoagulants, as well as to review the recommendation of international guidelines for the management of anticoagulants in urgent and elective endoscopic procedures.

**INTRODUCTION**

Endoscopic procedures are commonly performed in patients taking antiplatelet and anticoagulants agents. These antithrombotic medications reduce thromboembolic events by inhibiting platelet aggregation and coagulation and are the most widely prescribed agents in both primary and secondary care in many patients[1]. In addition, a growing number of subjects have an indication for combination therapy which increases the overall risk of bleeding, in particular that from the gastrointestinal tract[2].

The aim of this review is to analyze current evidence from literature assessing, for each procedure, the basal bleeding risk and the risk of bleeding during anticoagulant therapy, as well as to review the recommendation of American society for gastrointestinal endoscopy (ASGE)[3], European society of gastrointestinal endoscopy and British society of gastroenterology (ESGE/BSG)[4] and Asian pacific association of gastroenterology and Asian pacific society for digestive endoscopy (APAGE/APSDE)[5] for the management of anticoagulants in urgent and elective endoscopic procedures.

**ANTICOAGULANT THERAPIES**

Antithrombotic therapies may be classified in antiplatelet agents (APA) and anticoagulants.

***Antiplatelets***

APA interfere in specific steps of platelets activation process and include several agents namely aspirin [acetylsalicylic acid (ASA)], nonsteroidal anti-inflammatory drugs (NSAIDs), *P2Y12* platelet receptor blockers and other agents such as glycoprotein *IIb/IIIa* antibodies and receptor antagonists and the competitive and selective inhibitors of protease-activated receptor-1.

ASA is used to inhibit platelet aggregation by irreversibly blocking the ciclooxygenase pathway, resulting in the suppression of prostaglandin and thromboxane biosynthesis from arachidonic acid in platelets. After cessation of ASA, 7 to 9 d are required to full recover the platelet function. Dipyridamole reversibly prevents platelet activation by multiple mechanisms such as the inhibitions of cyclic nucleotide phosphodiesterase and the blocking of the uptake of adenosine. Dipyridamole has an elimination half-life of 12 h and a duration of action of about two days after discontinuation. Thienopyridine agents represent the most common antiplatelet drugs used following ASA. These agents selectively inhibit adenosine diphosphate-induced platelet aggregation, with no direct effects on the metabolism of arachidonic acid[6]. The class of *P2Y12* platelet receptor blockers is broad and include ticlopidine, clopidogrel and the most recent third-generation thienopyridine agents such as prasugrel and ticagrelor. Ticlopidine, clopidogrel and prasugrel are prodrugs which achieve their antiplatelet effects through active metabolites irreversibly inactivating the *P2Y12* receptor. The hepatic metabolism of prasugrel is more rapid (occurs in a single hepatic step) and less influenced by cytochrome *P450* polymorphisms than clopidogrel[7,8]. Prasugrel and ticagrelor induce a more rapid and pronounced inhibition of platelet aggregation compared to clopidogrel. Unlike the other thienopyridine which requires discontinuation of at least 5 to 7 d to recover adequate platelet function, ticagrelor induce reversible inhibition of the *P2Y12* receptor which permits a shorter interval of interruption (3 to 5 d)[9]. Other antiplatelet medication include the antagonists of the platelet glycoprotein (GP)*IIb/IIIa* (*e.g* abciximab, roxifiban, eptifibatide, tirofiban, orbofiban, sibrafiban) which play their antiplatelet effects by blocking the final common pathway of platelet aggregation. Current guidelines recommend antiplatelet agents for the secondary prevention of cardiovascular disease[10]. Conversely, these drugs are not recommended for the primary prevention of cardiovascular disease, although some evidence weakly supports the advantage of aspirin in patients with hypertension and impaired renal function or who are at high cardiovascular risk (10-year risk > 20%)[11,12].

***Anticoagulants***

Anticoagulants are prescribed in several clinical setting such as the prevention of stroke in patients with atrial fibrillation and as prophylaxis and treatment of venous thromboembolism. These drugs prevent the clotting of blood by inhibiting one or more steps in the coagulation cascade through several mechanisms of actions including both direct and indirect enzymatic blocking, the antagonism of vitamin K–dependent clotting factors and the binding to antithrombin. Available drugs include unfractionated heparin, low molecular weight heparins, fondaparinux, vitamin K antagonists (*e.g* warfarin), direct factor *Xa* inhibitors (*e.g* rivaroxaban, apixaban, edoxaban) and direct thrombin inhibitors which prevent thrombin from cleaving fibrinogen to fibrin and include both parenteral agents (bivalirudin, argatroban and desirudin) and oral agent (dabigatran etexilate). The efficacy and the bleeding risk related to the use of anticoagulants depends on the drug used and the clinical setting[13].

Data from the pivotal clinical trials (RE-LY, ROCKET AF, ARISTOTLE, and ENGAGE AF-TIMI 48) suggest that direct oral anticoagulants (DOACs) are not inferior to warfarin for the prevention of stroke or systemic embolism in subjects with non-valvular atrial fibrillation and that these drugs also had significant reductions in hemorrhagic stroke and intracranial hemorrhage compared to warfarin which results in a lower risk of stroke and mortality[14]. Furthermore, DOACs were associated with less severe major bleedings than those related to warfarin. Nevertheless, the rates of gastrointestinal (GI) bleeding are increased in non-valvular atrial fibrillation patients treated with DOACs and in particular with dabigatran 150 mg bid, rivaroxaban, and edoxaban but it seems to occur the least in patients receiving apixaban compared with vitamin K antagonists therapy[15]. In this regard, Abraham et al reported that the rates of GI bleeding were significantly increase with rivaroxaban than dabigatran (HR 1.20; 95%CI: 1.00-1.45), whereas apixaban was associated to a lower risk of GI bleeding than dabigatran (HR 0.39; 95%CI: 0.27-0.58; *P* < 0.001) or rivaroxaban (HR 0.33; 95%CI: 0.22-0.49; *P* < 0.001)[16]. However, despite the risk of bleeding associated with DOACs, this risk is counterbalanced by the effectiveness of these drugs in preventing the risk of stroke that is considered by patients as a “fate worse than death"[17]. A specific antidote is available for dabigatran (idarucizumab)[18], but not for the others DOACs. Recently, a direct factor *Xa* inhibitor has showed promising results against rivaroxaban, apixaban and edoxaban (andexanet alfa)[19].

**RISK OF BLEEDING ASSOCIATED WITH ENDOSCOPIC PROCEDURES**

The risk of bleeding associated with endoscopic procedures is variable and mainly depends on the type of procedure performed (diagnostic, low-risk or high-risk operative), on the type of antiplatelet or anticoagulant therapy and on patient’s comorbidities. In the following paragraphs, we will review and discuss, for each procedure, the evidence from literature assessing the risk of basal bleeding and the risk of bleeding in patients taking anticoagulant therapies.

***Diagnostic endoscopy***

Diagnostic procedures as esophagogastroduodenoscopy (EGD), colonoscopy or sigmoidoscopy, including mucosal biopsy present a low risk of procedural bleeding, as showed by several studies[20-21].

Similarly, the risk of bleeding during diagnostic endoscopy is not increased in patient assuming ASA, clopidogrel or warfarin. In this regard, a prospective, single-blind, randomized study in healthy volunteers, showed that on a sample of 405 antral biopsies and 225 duodenal biopsies performed during 90 EGD in 45 subjects receiving aspirin or clopidogrel, no bleeding events were noted in the clopidogrel group after 350 biopsies, while in the aspirin group, only one minor endoscopic bleeding event was reported, in the absence of clinical events[22].

Moreover, a prospective single-arm study including 112 Japanese outpatients receiving antithrombotic agents showed that after 101 biopsies performed during EGD or colonoscopy, no patients complained of any bleeding symptoms in the 2 wk observation period after biopsy. In addition, authors didn’t find significant differences between patients receiving single and multiple antithrombotic agents, as well as between patients not receiving and receiving warfarin[23].

Conversely, no data is currently available on the risk of bleeding after biopsies in patients taking DOACs.

A retrospective multicenter study of double balloon enteroscopy complications performed in the United States reported a risk of gastrointestinal bleeding of 0.2% associated with the procedure[24]. Unfortunately, no studies assessing the risk of bleeding during double balloon enteroscopy in patients on antiaggregant or anticoagulant therapy have been performed up to date. In the absence of data the risk is uncertain, but it can be assimilated however to that of other endoscopic diagnostic procedures.

***Polypectomy***

Polypectomy may be complicated by intraprocedural or immediate bleeding (IPB) and by post-procedural bleeding (PPB). While IPB is often self-limited and may be however controlled during the procedure, conversely, PPB may be worrisome, since it arises after the procedure when the patient has already been discharged.

Data from large series show a PPB ranging from 0.07% to 2.2%[25-28]. A large report from the English National Health Service Bowel Cancer Screening Programme shows an overall bleeding rate of 0.65 %, a rate of severe bleeding requiring transfusion of 0.04 %, and an increased bleeding risk attributable to polypectomy of 11.14-fold[29].

Exploring factors associated with portal pressure gradient, a prospective, cross-sectional study of 5152 patients undergoing polypectomy, showed that age ≥ 65 years, cardiovascular or chronic renal disease, anticoagulant use, polyp size > 1 cm, pedunculated polyp or laterally spreading tumor, poor bowel preparation and use of pure cutting current were significant risk factors for PPB[30]. Another case-control study confirmed that polyp size were associated with portal pressure gradient, with an increased risk of hemorrhage of 9% for every 1 mm increase in polyp diameter (OR 1.09; 95 %CI 1.0-1.2; *P* = 0.008)[31].

Post-polypectomy bleeding in patients on antithrombotic therapy has a different impact.

The risk in patients on ASA or NSAID was generally judged to be low.

One of the first studies performed on this field, showed as the risk of bleeding did not seem to be affected by NSAID use. In fact, although the use of NSAIDs increased the incidence of minor self-limited bleeding, an increase in the rate of major bleeding was not observed[32]. In addition, a revision of a cohort of 5593 patients and 1657 polypectomies clearly showed that the use of antiplatelet agents during polypectomy was not associated with an increase in post-polypectomy bleeding[33].

In a prospective multicenter trial, including a total of 1015 polyps <  10 mm removed by cold snare in 823 patients, 15% of them taking low dose aspirin or ticlopidine, reported a higher immediate post-polypectomy bleeding in patients taking APA (6.2 % *vs* 1.4 %; *P* < 0.001). Nevertheless, all bleeding episodes were intraprocedural and successfully treated, while no delayed PPB occurred[34].

Although low in patients receiving aspirin or NSAID, the risk of post-polypectomy bleeding is higher in patients receiving thienopyridine or warfarin.

Despite some studies advocate a low risk of post-polypectomy bleeding in patients taking thienopyridine[35], a meta-analysis of 5 observational studies including 574 subjects on clopidogrel therapy and 6169 controls showed an overall higher risk of PPB on continued clopidogrel (RR 2.54, 95%CI: 1.68-3.84, *P* < 0.00001), with a non-significant risk of immediate bleeding (RR of 1.76, 95%CI: 0.90-3.46, *P* = 0.10) and a significant higher risk of delayed bleeding (RR of 4.66, 95%CI: 2.37-9.17, *P* < 0.00001)[36].

While patients on anticoagulation therapy may safely undergo colonoscopy, current practice guidelines consider polypectomy a high-risk procedure for which anticoagulation must temporarily be discontinued. Despite this, this estimate of risk of bleeding is difficult to quantify with accuracy, since it depends on many variables, especially by the size of the polyp to be removed and International Normalized Ratio (INR) values.

A small single-center retrospective study performed on 21 patients receiving long-term anticoagulation with warfarin with an average INR of 2.3 and undergoing polypectomy, reported no episodes of PPB[37].

Another retrospective study supported the safety of polypectomy of small polyps < 10 mm without interruption of anticoagulation showing that, in a sample of 225 polypectomies with subsequent prophylactic placement of hemoclips, the rate of severe PPB requiring transfusion and of minor PPB not requiring treatment were 0.8% and 1.6%, respectively[38].

Similarly, a Japanese prospective randomized controlled trial of 70 patients randomized to cold snare or traditional polypectomy of lesions up to 10 mm without interruption of warfarin, confirmed that the incidence of PPB after cold snare resection was acceptable and that this technique reported a lower incidence of immediate (5.7% *vs* 23.0%) and delayed bleeding (0% *vs* 14%) compared to traditional polypectomy, even without interruption of anticoagulant therapy[39]. Despite this, further data are needed in order to better assess the setting in which a polypectomy on warfarin therapy could be performed safely.

To date, current guidelines recommend discontinuing warfarin 5 d before high-risk endoscopic procedures in patients at low thrombotic risk and discontinuing warfarin 5 d before high-risk endoscopic procedures with low molecular weight heparin (LMWH) bridging in patients at high thrombotic risk. In this regard, also the role of bridging need to be better assessed, as recent data suggest that patients undergoing bridging with LMWH are at higher risk of procedural-related bleeding compared to patients than those not undergoing bridging with LMWH or continuing warfarin therapy[40,41]. On the same line, a recent study showed as patients discontinuing anticoagulant with LMWH bridging, as suggested by guidelines, had a higher PPB rate compared to patients continuing anticoagulants (10.8% *vs* 19.6%, *P* = 0.087)[42]. Based on these observations, bridging recommendation should be revised, and the choice should be individualized taking into account the hemorrhagic and thromboembolic risk of the individual patient.

With regard to DOAC, the risk of PPB is not well known. The aforementioned study, comparing post-polypectomy complication rates in 218 patients receiving oral anticoagulants (73 DOACs, 145 warfarin) and 218 patients not receiving anticoagulant therapy, showed that the PPB was similar between DOAC and warfarin and higher for both compared with controls (13.7% *vs* 13.7% *vs* 0.9%, *P* < 0.001)[42].

Similarly, the second mentioned analysis of 11504 comparing patients on antithrombotic therapy (1590 DOAC, 3471 warfarin, and 6443 clopidogrel) and 599983 control undergoing colonoscopy with polypectomy or endoscopic mucosal resection, showed that subjects undergoing DOACs did not have a statistically significant increased risk gastrointestinal bleeding, as well as cerebrovascular accident or myocardial infarction and hospital admissions compared with controls. On the contrary, clopidogrel and warfarin were associated with increased odds of PPB, cerebrovascular accident or myocardial infarction and hospital admissions compared with controls[43].

***Endoscopic mucosal resection***

The overall risk of intraprocedural and delayed bleeding after Endoscopic mucosal resection (EMR) has been estimate between 3.7%-11.3 % and 0.6%-6.2 %, respectively, and it is therefore higher compared with polypectomy[4].

The risk of bleeding after EMR is associated with location and the size of the lesion. Esophageal EMR has a greater risk of bleeding ranging from 4% to 20%[44-47]. Moreover a study showed as esophageal EMR (OR 2.5, 95%CI: 1.2-5, *P* = 0.0009) and lesion size (OR 1.24, 95%CI: 1.1–1.5, *P* = 0.003) were independently associated with higher risk of early bleeding in EMR, when controlled for age, gender and NSAIDs or clopidogrel therapy[48]. Besides, duodenal EMR presents a risk of delayed bleeding between 6.3 and 12.3[47,48]. The risk of hemorrhages seems to be lower for EMR of lesions smaller than 1 cm[49].

Recently, a delayed bleeding prediction model (GSEED-RE2) that takes into account several variables including lesion size, proximal location, comorbidity, and antiplatelet or anticoagulant therapy, has been proposed showing higher values of area under the curve (0.69-0.73; 95%CI: 0.59-0.80) compared to previous models[50].

Prophylactic measures, may help to reduce the risk of delayed bleeding after EMR, for instance with the placement of hemoclips. In this regard, a retrospective study of 524 lesions 2 cm or larger resected by EMR showed that the delayed bleeding rate was 1.8% in the clipped group *vs* 9.7% in the not clipped group, and that the absence of clipping (OR 6.0; 95%CI: 2.0-18.5) was independently associated with delayed hemorrhage[51]. On the contrary other two studies did not find any significant difference in the rate of PPB between patients with and without prophylactic placement of hemoclips[52,53].

With regard to the risk of bleeding associated with anticoagulant therapy, data from two large prospective intention-to-treat studies of 302 EMR for colonic laterally spreading tumors ≥ 20 mm performed in 288 patients, showed that use of aspirin (OR 6.3, *P* = 0.005), was independently associated with risk of bleeding at multivariate analysis[54]. On the contrary, temporary discontinuation of anticoagulants seems to be safe. A retrospective study on a cohort of 798 patients undergoing 1716 EMR, all of them stopping antiplatelets and anticoagulants 7 d before EMR and resuming clopidogrel 2 d after EMR, showed that the temporary cessation of clopidogrel before EMR and its prompt resumption was not associated with an increased risk of gastrointestinal bleeding[55]. No data is available for other anticoagulants, including DOACs, for the risk of bleeding after EMR.

Based on these data, EMR is therefore considered a high risk endoscopic procedure for management of anticoagulant therapy.

***Endoscopic submucosal dissection***

Endoscopic submucosal dissection (ESD) is a more recent and more complex technique, with a growing popularity worldwide. Although a more radical resection, the risk of IPB and of PPB seems to be higher compared to EMR. A meta-analysis of 15 nonrandomized studies comparing ESD with EMR confirmed higher ‘en bloc’ resection rates (OR 13.87, 95%CI: 10.12-18.99) and higher curative resection rates for ESD compared to EMR (OR 3.53, 95%CI: 2.57-4.84), with the disadvantage of an higher procedure-related bleeding (OR 2.20, 95%CI: 1.58)[56].

Nevertheless, the risk is lower for esophageal and colonic ESD, and higher for gastric ESD. A meta-analysis of 15 studies with a total of 776 ESD procedures performed for resection of esophageal neoplasia showed a pooled estimate of PPB of 2.1 % (95%CI: 1.2-3.8)[57]. Similarly, in another meta-analysis of 22 studies with a total of 2841 colonic ESD the pooled estimate of PPB was 2.0 % (95%CI: 1.0-2.0)[58].

On the contrary, gastric ESD present a risk of PPB ranging from 3.6% to 6.9% as reported in 7 studies including > 11000 procedures[59-65].

With regard to the risk of bleeding associated with anticoagulant therapy, most of the data come from gastric setting. In patients who do not discontinue aspirin before gastric ESD, aspirin use is independently associated with PPB (RR 4.49; 95%CI: 1.09-18.38)[66]. The risk of bleeding in patients who discontinue aspirin before gastric ESD is controversial. Some studies show that the risk of bleeding is increased even after its temporary withdrawal[67,68], while other studies showed that its discontinuation is not independently associated with delayed bleeding after the procedure[69,70]. Similarly, in colorectal ESD, discontinuation of antithrombotic therapy is not associated with post-procedural bleeding[71,72]. Beyond aspirin, no consistent data is available on the effect of other anticoagulants, such as clopidogrel, ticagrelor and DOACs, on the risk of bleeding associated to ESD.

***Endoscopic dilatation***

According to current literature data, no significant risk of bleeding is reported neither in esophageal dilations[73-77], nor in those of the colon[78-84]. Beside this, no study evaluated so far the risk of bleeding due to endoscopic dilatation in patients under APA or anticoagulant therapy.

***Endoscopic stenting***

The risk of bleeding after endoscopic stenting is difficult to assess precisely, due to heterogeneity of the type of stent, the anatomical site and indication (benign *vs* malignant), and it still controversial.

With regard to esophageal stenting, several studies report a risk of bleeding ranging between 1% and 8%[85,86].

Besides, in the setting of gastroduodenal stenting, a systematic review of 32 case series of stent placement in 606 patients with malignant symptomatic gastroduodenal obstruction showed an incidence of bleeding of 0.5%[87].

Moreover, two systematic reviews assessed the incidence of bleeding in colonic stenting. The first one, evaluating 29 case series and 598 stent placements, showed a 4.5% bleeding rate[88]. The second, including 54 non-randomized studies with the use of stents in a total of 1198 patients, did not report any case of bleeding[89].

Also in this case, no study evaluated so far the risk of bleeding due to endoscopic stenting in patients taking APA or anticoagulants.

***Percutaneous endoscopic gastrostomy***

Percutaneous endoscopic gastrostomy (PEG) is performed through the perforation of the abdominal and the anterior stomach walls. Therefore, bleeding from some vessel placed in the path of the puncture or, rarely, a hematoma of the wall, may occur. A recent multicenter prospective cohort study including 950 patients undergoing PEG placement or replacement, showed a 1% bleeding rate[90].

In another retrospective, single-center study of 990 patients undergoing PEG placement, the incidence of bleeding was 1.6%, and multivariate analysis demonstrated no association between periprocedural use of aspirin (at any dose) or clopidogrel and post-PEG bleeding[91].

Also in this setting, no data is available on the risk of bleeding after PEG placement in patients taking prasugrel, ticagrelor or DOACs.

***Esophageal variceal ligation***

Band ligation of esophageal varices is generally performed as emergency treatment for active upper variceal bleeding and, in this setting, measures for management of anticoagulants in urgent endoscopy should be applied. The band ligation can be also performed as an elective procedure for primary prophylaxis of varices that have never previously bleed. A case-control study showed an incidence of delayed bleeding around 3.4%. In the same study, aspirin or anticoagulation were not independently associated with the risk of bleeding, even though they were taken by a small minority (1.3%) of patients[92].

No data is available regarding the risk of bleeding in patients taking other anticoagulants, such as thienopyridines, ticagrelor or DOACs.

***Endoscopic retrograde cholangiopancreatography***

The risk of bleeding during endoscopic retrograde cholangiopancreatography (ERCP) is reported in about 0.1% to 2% of procedures with sphincterotomy[93].

In this regard, a meta-analysis showed that, even if sphincterotomy before stent placement reduces the risk of post ERCP pancreatitis (OR = 0.34, 95%CI: 0.12-0.93, *P* = 0.04), it is associated with a higher risk of post-ERCP bleeding (OR = 9.70, 95%CI: 1.21-77.75, *P* = 0.03)[94].

Patient factors affecting the risk of bleeding after sphincterotomy include anticoagulant therapy, presence of coagulopathies, and active cholangitis.

On the other side, procedural factors affecting the risk of bleeding are the operator’s experience and the type of current used during the sphincterotomy. As a matter of fact, a meta-analysis confirmed as a mixed-current was associated with a lower risk of bleeding (12.2%-95%CI, 4.1%, 20.3%) compared to pure-cut current (37.3%-95%CI, 27.3%, 47.3%), with a similar risk of pancreatitis[95].

Another meta-analysis showed that, compared to sphincterotomy, endoscopic papillary balloon dilatation is associated with a lower risk of bleeding (OR = 0.12, 95%CI: 0.04-0.34, *P* < 0.01), even if presented a significant higher incidence of pancreatitis (OR = 2.79, 95%CI: 1.74-4.45, *P* < 0.0001)[96].

The risk of bleeding after sphincterotomy in patients taking antiplatelet agents is not completely defined. Several studies did not find any significant additional risk of bleeding in patients taking antiplatelet agents[93, 97-100]. Beside this, a retrospective study showed an higher incidence of post-sphincterotomy bleeding in patients continuing aspirin until the day of sphincterotomy, compared to patients that had never taken aspirin (9.7% *vs* 3.9%, *P* = 0.01)[101].

No data from literature is currently available regarding risk of bleeding after biliary lithotripsy and cholangioscopy in patients taking APA or anticoagulants.

***Endoscopic ultrasound with fine-needle aspiration***

Endoscopic ultrasound (EUS) plus fine-needle aspiration (FNA) is performed for diagnosis and locoregional staging of esophageal, gastric, rectal, and pancreatic cancers. Despite the procedure include the puncture of the tissue, the reported incidence of bleeding is low.

A large systematic review of 51 articles with a total of 10941 patients undergoing endoscopic ultrasound with fine-needle aspiration (EUS-FNA), showed a bleeding in 0.17% of procedures, and most bleeding complications occurred after FNA performed on pancreatic lesions[102]. Moreover, a prospective controlled study assessed the incidence of bleeding after 222 EUS-FNA procedures in patients undergoing aspirin/NSAIDs and LMWH compared to patients didn't take these therapies. Bleeding occurred in 0% (0/26), 33.3% (2/6) and 3.7% (7/190) of the patients in the aspirin/NSAIDs, LMWH, and control groups, respectively (*P* = 0.023)[103]. These data confirm as EUS-FNA is a safe procedure, even if performed in patients taking aspirin or NSAIDs.

No data is available regarding the risk of bleeding after EUS-FNA in patients taking anticoagulants.

***Peroral endoscopic myotomy***

Peroral endoscopic myotomy is a new endoscopic technique for the treatment of esophageal achalasia, with excellent results[104].

Since the view on the vasculature around the muscle fibers is optimal and only few vessels are encountered in the submucosal tunnel, major bleeding is not frequent during the procedure. According to a recent meta-analysis massive hemorrhage was reported in 0.2% of procedures[105].

Currently, no data on the risk of bleeding in patients taking antiplatelets or anticoagulants is available.

**MANAGEMENT OF ANTICOAGULANTS IN ELECTIVE PROCEDURES**

Currently, international practice guidelines, provide specific recommendation on management of anticoagulant therapy in the periendoscopic period.

The management of anticoagulants depends on the type of molecule, the estimated risk of bleeding due to the endoscopic procedure, and the underlying thrombotic risk deriving from cardiovascular disease.

With regard to the first parameter, ASGE, ESGE/BSG and APAGE/APSDE guidelines identifies specific risk groups (Table 1). The low-risk group includes all endoscopic diagnostic procedures (including EGD, colonscopy and enteroscopy with or without mucosal biopsies, EUS without FNA) and ERCP. The high-risk group includes polypectomy, EMR, ESD, therapeutic enteroscopy, ERCP with biliary or pancreatic sphincterotomy, ampullectomy, EUS with FNA, dilatation of strictures and PEG placement. These two groups are homogeneous between the guidelines, with the exception of enteral stents that are still controversial and classified as low-risk by ASGE and APAGE/APSDE, and as high-risk by ESGE/BSG. Peroral endoscopic myotomy is not classified by any of the guidelines but could be ascribed to high-risk procedures. Moreover, APAGE/APSDE consider a third group of ultra-high risk procedures including ESD and EMR of polyps > 2 cm, for which specific indication are provided.

On the other hand, the assessment of cardiovascular risk differs between the three guidelines and is summarized in Table 2.

***Aspirin and NSAIDs***

With regards to aspirin, ASGE guidelines judge the use of low doses of aspirin and NSAIDs safe and suggest to continue these drugs in the periendoscopic period[3].

On the other side, ESGE/BSG and APAGE/APSDE guidelines recommend continuing aspirin therapy for almost all endoscopic procedures, with some different exceptions.

ESGE/BSG recommend aspirin discontinuation, on an individual basis, in patients undergoing ESD, EMR for upper gastrointestinal lesions and colonic lesions > 2 cm, and ampullectomy[4]. Beside this, APAGE/APSDE recommend discontinuation of aspirin in all patients undergoing ESD and EMR of all large (> 2 cm) polyps[5] (Figure 1).

***Thienopyridines******(clopidogrel, prasugrel, ticagrelor)***

Recommendations on thienopyridines management are provided depending on the estimate procedure risk: (1) For patients undergoing low-risk endoscopic procedures, all guidelines recommend to continue thienopyridine therapy. With regard to dual antiplatelet therapy, ESGE suggest to continue dual antiplatelet therapy, while APAGE/APSDE advice to don’t stop both antiplatelet agents; (2) For patients undergoing high-risk endoscopic procedures, ASGE and ESGE/BSG recommend to assess before the cardiovascular risk (CVR, Table 2): If low CVR, stop thienopyridine before endoscopy. In the case of dual APA therapy, both ASGE and ESGE/BSG agree to continue aspirin if already prescribed; if high CVR, ASGE recommend to discontinue thienopyridine at least 5 d before or switch to ASA, while ESGE/BSG suggest to consider discontinuation of thienopyridine 5 d before only after 12 mo following insertion of drug-eluting coronary stent or after 1 mo suggest insertion of bare metal coronary stent.

APAGE/APSDE recommends to discontinue thienopyridine at least 5 d before endoscopy in all patient undergoing a high-risk endoscopy procedure regardless of the CVR.

With regard to patients with dual APA therapy, all guidelines agree to withhold the thienopyridine and continue aspirin. Moreover, according to APAGE/APSDE ultra-high risk procedures may require stopping both antiplatelet agents (Figure 2).

After the procedure, the suggested management differs according to the guidelines: ASGE suggest to resume thienopyridine after the procedure once hemostasis is achieved; in this setting, a loading dose of thienopyridine should be considered among patients at risk for thrombosis[3]; ESGE/BSG recommend that thienopyridine should be resumed up to 48 h after the procedure depending on the perceived bleeding and thrombotic risks[4]; APAGE/APSDE recommend early resumption of thienopyridine within 5 d after endoscopic hemostasis in patients with drug-eluting coronary stents[5].

***Warfarin***

Also for warfarin, recommendations are provided depending on the estimate procedure risk: (1) For patients undergoing low-risk endoscopic procedures, all guidelines recommend to continue warfarin 5 d before the procedure. In addition ESGE/BSG and APAGE/APSDE suggest to check the INR the week before endoscopy to ensure it is in the normal range[4,5]; (2) For patients undergoing high or ultra-risk endoscopic procedures, all guidelines recommend to assess before the CVR (Table 2): If low CVR, discontinue warfarin 5 d before the procedure and check the INR before the procedure to ensure values < 1.5 according to ESGE/BSG[4] or less conservative values < 2 according to APAGE/APSDE[5]; if high CVR, discontinue warfarin 5 d before the procedure ads administer bridge therapy with LMWH. In addition, ESGE/BSG recommend to withdraw last dose of LMWH more than 24 h before the procedure[4] (Figure 3).

All guidelines recommend resuming warfarin the same day/evening of the procedure after a proper hemostasis has been achieved. Moreover, for patients with high CVR, ESGE/BSG and APAGE/APSDE specify to continue LMWH until therapeutic INR range has been achieved.

***DOACs (dabigatran, rivaroxaban, apixaban, edoxaban)***

For patients undergoing low-risk endoscopic procedures ASGE and APAGE/APSDE recommend to continue DOAC in the periendoscopic period, similarly to what is suggested for warfarin[3,5]. On the contrary, ESGE/BSG guidelines recommend omitting the DOAC dose the morning of procedure. Nevertheless, this last statement is reported as a weak recommendation and supported by very low quality evidence[4].

For patients undergoing high-risk endoscopic procedures, all guidelines recommend to discontinue DOACs before endoscopy for the appropriate drug-specific interval, adjusting for creatinine clearance[3-5]. In this regard, ESGE/BSG and APAGE/APSDE specify to take last dose of DOAC ≥ 48 h before procedure[4,5] (Figure 4).

Unlike reintroduction of warfarin, which results in delayed anticoagulation for several days, a therapeutic intensity of anticoagulation is restored within 3 h of taking a therapeutic dose of a DOAC. Because of the high risk of bleeding associated with the therapeutic intensity of anticoagulation after an invasive procedure, guidelines suggest a delay in reintroducing DOACs after a high-risk procedure. This delay will depend on the bleeding risk associated with the procedure and will usually be 24-48 h [4]. For procedures with a significant risk of delayed bleeding such as EMR or ESD, a longer period of discontinuation may be considered for patients with a relatively low-thrombotic risk.

According to ASGE guidelines, if DOACs cannot be restarted within 24 h after a high-risk procedure because of concern regarding the adequate hemostasis due to their short onset of action, then thromboprophylaxis (*e.g.,* LMWH bridge) should be considered for patients at high risk for thromboembolism[3].

On the contrary, APAGE/APSDE recommends early resumption of DOACs soon after the procedure after adequate hemostasis has been achieved.

**MANAGEMENT OF ANTICOAGULANTS IN URGENT PROCEDURES**

***Antiplatelets agents***

ASGE guidelines recommend consultation with the prescribing specialist before stopping APAs during gastrointestinal bleeding in patients for which the cardiovascular risk overcome the potential consequences of bleeding, namely patients with: (1) Placed drug eluting intracoronary stents within 1 year, (2) Insertion of a bare metal intracoronary stent within 30 d, or after an acute coronary syndrome within 90 d[3].

The ESGE/BSG guidelines propose an algorithm in which patient management depend form primary prophylaxis or secondary prophylaxis.

The algorithm recommend to withholding aspirin until the third day after endoscopic treatment of high-risk stigmata in patient taking APA for secondary prophylaxis[4].

In this regards two studies have shown how for patients taking low-dose aspirin for secondary cardiovascular prophylaxis the all-cause mortality was lower if aspirin was continued[106,107].

Beside this, APAGE/APSDE guidelines recommend to withhold aspirin only in patients with serious or life-threatening bleeding in places where endoscopy is not readily available[5].

With regard to patients taking dual antiplatelet therapy having an acute gastrointestinal bleeding, both ESGE/BSG and APAGE/APSDE recommend continuation of aspirin and withholding clopidogrel. In these patients, resumption of therapy preferably within 5 d after endoscopic hemostasis is recommended by APAGE/APSDE guideline, whereas ESGE/BSG and ASGE suggest consultation with a cardiologist for the management after urgent endoscopic treatment.

***Warfarin***

The estimated incidence of gastrointestinal bleeding in patients in anticoagulant therapy ranges between 1%-4% per year[108].

The management of patients using anticoagulants with active gastrointestinal bleeding has been the focus of numerous studies for many years.

The ESGE/BSG and ASGE guidelines recommend to stop the therapy and correct the INR in patients taking vitamin K antagonists with signs of severe bleeding, before performing urgent endoscopy[109-111]. Moreover it also recommend to not delay endoscopy if INR < 2.5. In this regard,Choudari and colleagues described a retrospective series of 52 patients which developed bleeding while taking warfarin. The outcome and the endoscopy efficacy in patients with an INR corrected between 1.5 and 2.5 were good and similar to those recorded in patients not taking anticoagulants. Therefore, even a partial correction of the INR seems to be associated with a good outcome[112].

In a retrospective cohort study of 233 consecutive anticoagulated patients undergoing urgent endoscopy with successful haemostasis, 95% of them had an INR between 1.3 and 2.7. The rate of re-bleeding was 23% but INR was not a predictor of re-bleeding on multivariable analyses[113].

Another retrospective study of patients taking warfarin at the moment of admission for gastrointestinal bleeding, reports 55 patients with INR value of 4.0 or greater (supratherapeutic) and 43 with INR in the range 2.0 to 3.9. Patients with supratherapeutic INRs were more likely to have a gastrointestinal pathology supporting a role for endoscopic evaluation of these patients but no differences in rete of re-bleeding were recorded[114].

A large systematic review of 1869 patients with nonvariceal upper GI bleeding, showed that INR value at initial presentation of the event was found not to predict the risk of rebleeding. Rather, the finding of values of INR > 1.5 during non-variceal digestive haemorrhage was associated with an increase in mortality after correction in patients with specific comorbidities. So in this study the INR value was found to be useful in the risk stratification than as a predictor of rebleeding[115]. Furthermore, in a retrospective case-control study Irwin and colleagues have highlighted that patients with supratherapeutic INR at the time of gastrointestinal bleeding had a lower mortality rate 30 d after the event compared to patients not taking warfarin[116].

These evidences suggest that in the clinical practice could not be so useful to normalize the INR in all patients delaying the timing of endoscopy. Hence, as the ASGE and ESGE/BSG practice guidelines suggest, the endoscopic treatment can be considered effective and relatively safe in patients with INR values ​​< 2.5[3,4].

In the management of a patient with active gastrointestinal bleeding, the decision to correct coagulopathy (stopping, reducing or reversing it) it’s often difficult because it concerns patients with a risk of thromboembolic consequences.

The thrombotic risk after transient withdrawal of anticoagulant therapy, in acute gastrointestinal bleeding settings, was explored by 2 small studies conducted on 27 and 28 patients. The withdrawal time was variable between 5 and 14 d, in one of the two studies it was also administered vitamin K or frozen plasma was given to 7 patients to reverse anticoagulation. Both studies showed a low risk of thrombotic complications[117,118].

All the guidelines (ASGE, ESGE/BSG and APAGE/APSDE) agree and advice on the urgent reversal in all patients presenting with a life-threatening gastrointestinal bleeding, regardless of therapeutic or supra-therapeutic INR elevations[3-5, 119].

Specifically, ESGE/BSG recommends estimating the cardiovascular risk of the patient in consultation with a cardiologist before starting INR correction[4].

For patients without signs of active bleeding and hemodynamically stable the usefulness of INR correction should be evaluated on a case-by-case basis.

For an urgent correction of coagulopathy in patient taking warfarin, ASGE and ESGE/BSG guidelines suggest administration of prothrombin complex concentrates (PCC) or fresh frozen plasma (FFP)[3,4].

Moreover, ESGE/BSG highlight that it is preferable to use PCC in combination with intravenous vitamin K at the dosage of 5-10 mg K to prevent “rebound coagulopathy” limiting the use of FFP when PCC is not available[4].

Other guidelines, such as those of the APAGE/APSDE recommend only the combination of PCC concomitantly with low-dose vitamin K (< 5 mg instead of 5-10 mg) administration[5]. With regard to the latter, the decision to prefer low-dose vitamin K is based on evidence from four randomized clinical trials that the optimal dose of vitamin K to achieve a normalization of the INR value is between 1 and 2.5 mg[120-123].

Currently, there are no randomized clinical trials comparing prothrombin PCC and FFP for warfarin reversal in acute GI bleeding.

In 2014 Karaca *et al*[124] performed a prospective, nonrandomized, comparative study of 40 patients taking warfarin with upper gastrointestinal haemorrhage and INR > 2.1. Patients received either PCC or FFP. INR levels were reversed more quickly in PCC group at the second and sixth h (second h INR: 1.53 *vs* 4.50, *P* < 0.01, sixth h INR: 1.52 *vs* 2.41, *P* < 0.01). At time of endoscopy no patient in the PCC group had active bleeding compared with 7 in the FFP group (0% *vs* 35 %, *P* < 0.01).

Regarding the thrombotic risk, a meta-analysis of 2011 showed a similar risk for the two agents (approximates 1%)[125].

In addition to a faster onset of action, in the clinical practice PCC have other advantages, as no need for ABO matching, less risk for volume overload because of smaller transfusion volume and minimal risk of infectious transmission. On the contrary, a disadvantage is represented by the higher cost[126].

With regard to resumption of warfarin after bleeding event, APAGE-APSDE guidelines recommend to resume warfarin by day 3 after adequate haemostasis is achieved and to consider bridge with LMWH in patients with high thrombotic risk[5]. On the other side, ESGE/BSG suggest to resume warfarin between 7 and 15 d following the bleeding event if haemostasis is successfully achieved[4].

***DOACs***

The ESGE/BSG guidelines recommend the use of platelet transfusion or desmopressin for more severe bleeding although there is no clear evidence to support this approach[4]. As evidenced by recent data, platelet transfusion is associated with an increase in mortality[127].

The most recent endoscopic APAGE/APSDE guidelines, do not recommend making specific assays to estimate anticoagulant activity of DOACs in acute bleeding[5]. Prothrombin time and activated partial thromboplastin time could be inaccurate in DOACs activity evaluation and currently specific assays are not routinely available.

With the exception of patients with reduced renal clearance, generally, the best strategy in patients taking DOACs is their withdrawal due to their short half-lives.

Moreover, it should also be considered that, to date, antidotes and direct inhibitor of DOACs are available. In a multicenter open-label study of 503 patients, idarucizumab proved to be safe and effective in neutralizing the anticoagulant effect of dabigatran in emergency situations[15].

Another study assessed the effectiveness of andexanet alfa, a modified recombinant inactive form of human factor *Xa*, for bleeding associated with Factor *Xa* Inhibitors such as rivaroxaban, apixaban and edoxaban. The study showed that treatment with andexanet in patients with acute major bleeding associated with the use of a factor *Xa* inhibitor, markedly reduced anti-factor *Xa* activity, with an excellent or good hemostatic efficacy in 82% of patients[16].

After index bleeding episode, APAGE-APSDE recommend to resume DOACs by day 3 after haemostasis is successfully achieved, without heparin bridging[5].

On the contrary, ASGE and ESGE/BSG guidelines do not provide specific recommendation on resumption of DOACs[3,4].

**CONCLUSION**

Currently, there is sufficient evidence from the literature to assess the basal risk of bleeding associated to main endoscopic procedures in patients who do not take anticoagulants.

On the contrary, there are not sufficient data allowing to evaluate as this risk increases in patients taking anticoagulants, especially in some specific procedures (*e.g* endoscopic dilatation, stenting, esophageal variceal ligation, EUS-FNA *etc*). Nevertheless, the high risk associated with these procedures makes further studies unlikely to be carried out in the future.

To date, current international practice guidelines provide specific recommendations for the management of anticoagulants in endoscopy, with a good agreement and some major differences that have already been discussed (Figures 1-4).

Some of these differences may explained by a different proposed approach, different geographical area of reference and different year of publication.

Nevertheless, even with some discrepancy, guidelines only represent the guide for a common policy, over which a tailored approach should always be applied. On this line, a crucial point is the importance to assess the individual risk-benefit ratio.

As a matter of fact, interruption of anticoagulants has an undoubtful beneficial reducing the risk of bleeding during endoscopic procedures. Despite this, the thromboembolic risk that follows suspension of the therapy should not be underestimated. For this reason, the individual decision should be taken case by case taking into account every single factor.

In addition, since several endoscopic examination are commonly requested by general practitioner or by other consultants as “open access”, a preliminary gastroenterological visit with the patient is essential to better assess the procedure risk and the proper schedule for withdrawal of anticoagulants, as well as written information on anticoagulant resumption should be provided at discharge.

Moreover, when necessary, consultation with cardiologist or hematologist has a primary importance for a tailored approach on the single patient.

In the future, further studies are needed to better clarify the gray areas and topics on which the guidelines still present some controversial points. In the meantime, we suggest to strictly comply with international guidelines and to discuss difficult decisions within multidisciplinary boards.

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**Footnotes**

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**Table 1 Stratification of endoscopic procedures based on the risk of bleeding according to international guidelines**

|  |  |
| --- | --- |
| **Procedure risk group** | **Practice guidelines** |
| **ASGE[3]** | **ESGE/BSG[4]** | **APAGE/APSDE[5]** |
| Low-risk | (1) Diagnostic (EGD, colonoscopy, flexible sigmoidoscopy) including mucosal biopsy; (2) ERCP with stent (biliary or pancreatic) placement or papillary balloon dilation without sphincterotomy; (3) Push enteroscopy and diagnostic balloon-assisted enteroscopy; (4) Capsule endoscopy; (5) Enteral stent deployment (controversial); (6) EUS without FNA; (7) Argon plasma coagulation and (8) Barrett’s ablation  | (1) Diagnostic procedures +/– biopsy; (2) Biliary or pancreatic stenting; (3) Diagnostic EUS and (4) Device-assisted enteroscopy without polypectomy | (1) Diagnostic endoscopy with biopsy; (2) Endoscopic ultrasound without fine needle aspiration; (3) ERCP with biliary or pancreatic stenting; (4) Diagnostic push or device-assisted enteroscopy; (5) Video capsule endoscopy; (6) Oesophageal, enteral and colonic stenting and (7) Argon plasma coagulation  |
| High-risk | (1) Polypectomy; (2) Biliary or pancreatic sphincterotomy; (3) Treatment of varices; (4) PEG/PEJ placement; (5) Therapeutic balloon-assisted enteroscopy; (6) EUS with FNA; (7) Endoscopic hemostasis; (8) Tumor ablation; (9) Cystgastrostomy; (10) Ampullary resection; (11) EMR; (12) Endoscopic submucosal dissection and (13) Pneumatic or bougie dilation  | (1) Polypectomy; (2) ERCP with sphincterotomy; (3) Ampullectomy; (4) EMR; (5) ESD; (6) Dilation of strictures; (7) Therapy of varices; (8) PEG; (9) EUS with FNA and (10) Oesophageal, enteral or colonic stenting | (1) Polypectomy; (2) ERCP with sphincterotomy ± balloon sphincteroplasty; (3) Dilatation of strictures; (4) Injection or banding of varices; (5) PEG/PEJ placement; (6) EUS with FNA and (7) Ampullectomy  |
| Ultra-high-risk | NA | NA | (1) ESD; (2) EMR of large (> 2 cm) polyps  |

ASGE: American society for gastrointestinal endoscopy; ESGE/BSG: European society of gastrointestinal endoscopy and British society of gastroenterology; APAGE/APSDE: Asian pacific association of gastroenterology and Asian pacific society for digestive endoscopy; EGD: Esophagogastroduodenoscopy; EUS: Endoscopic ultrasound; FNA: Fine-needle aspiration; ERCP: Endoscopic retrograde cholangiopancreatography; PEG: Percutaneous endoscopic gastrostomy; PEJ: Percutaneous endoscopic jejunostomy; EMR: Endoscopic mucosal resection; ESD: Endoscopic submucosal dissection; NA: Not assessed.

**Table 2 Stratification of thrombotic risk according to international guidelines**

|  |  |
| --- | --- |
| **Thrombotic risk category** | **Practice guidelines** |
| **ASGE[3]** | **ESGE/BSG[4]** | **APAGE/APSDE[5]** |
| Low-risk | Anticoagulant therapy: (1) Bileaflet aortic valve prosthesis without AF and no other risk factors for CVA; (2) VTE > 12 mo previous and no other risk factors; (3) Atrial fibrillation with CHA2DS2-VASc score < 2 | Antithrombotic therapy:(1) Ischaemic heart disease without coronary stent; (2) Cerbrovascular disease; (3) Peripheral vascular disease. Anticoagulant therapy: (1) Prosthetic metal heart valve in aortic position; (2) Xenograft heart valve; (3) Atrial fibrillation without valvular disease; (4) > 3 mo after venous thromboembolism; (5) Thrombophilia syndromes  | Antithrombotic therapy: (1) Acute coronary syndrome or percutaneous coronary intervention > 6 mo ago; (2) Stable coronary artery disease. Anticoagulant therapy: (1) Non-valvular atrial fibrillation with a CHA2DS2-VASc score ≤ 5; (2) Prosthetic valve without atrial fibrillation; (3) > 3 mo after venous thromboembolism |
| Moderate-risk | Anticoagulant therapy: (1) Bileaflet aortic valve prosthesis and one or more of the following risk factors: AF, prior CVA or TIA, hypertension, diabetes, congestive heart failure, age > 75 yr; (2) VTE within the past 3-12 monon-severe thrombophilia (heterozygous factor V Leiden or prothrombin gene mutation);(3) Recurrent VTEactive cancer (treated within 6 mo or palliative)  |  |
| High-risk | Anticoagulant therapy: (1) Any mitral valve prosthesis; (2) Any caged-ball or tilting disc aortic valve prosthesis; (3) Recent (within 6 mo) CVA or TIA; (4) Atrial fibrillation with CHA2DS2-VASc score ≥ 2  | Antithrombotic therapy: (1) Drug eluting coronary artery stents within 12 mo of placement; (2) Bare metal coronary artery stents within 1 mo of placement. Anticoagulant therapy: (1) Prosthetic metal heart valve in mitral position; (2) Prosthetic heart valve and atrial fibrillation; (3) Atrial fibrillation and mitral stenosis; (4) < 3 mo after venous thromboembolism | Antithrombotic therapy: (1) Acute coronary syndrome or percutaneous coronary intervention 6 wk–6 mo. Anticoagulant therapy: (1) Non-valvular atrial fibrillation with a CHA2DS2-VASc score > 5; (2) Metallic mitral valve; (3) Prosthetic valve with atrial fibrillation; (4) < 3 mo after venous thromboembolism; (5) Severe thrombophilia (protein C or protein S deficiency and (6) antiphospholipid syndrome)  |
| Very high-risk |  |  | Antithrombotic therapy: Acute coronary syndrome or percutaneous coronary intervention < 6 wk |

ASGE: American society for gastrointestinal endoscopy; ESGE/BSG: European society of gastrointestinal endoscopy and British society of gastroenterology; APAGE/APSDE: Asian pacific association of gastroenterology and Asian pacific society for digestive endoscopy; AF: Atrial fibrillation; CVA: Cerebrovascular accident; VTE: Venous thromboembolism; TIA: Transient ischemic attack.



**Figure 1 Management of** **aspirin and nonsteroidal anti-inflammatory drugs in elective endoscopic procedures according to international guidelines.** NSAID: Nonsteroidal anti-inflammatory drugs; ASGE: American society for gastrointestinal endoscopy; ESGE/BSG: European society of gastrointestinal endoscopy and British society of gastroenterology; APAGE/APSDE: Asian pacific association of gastroenterology and Asian pacific society for digestive endoscopy; EMR: Endoscopic mucosal resection; ESD: Endoscopic submucosal dissection.



**Figure 2 Management of** **thienopyridines in elective endoscopic procedures according to international guidelines.** ASGE: American society for gastrointestinal endoscopy; ESGE/BSG: European society of gastrointestinal endoscopy and British society of gastroenterology; APAGE/APSDE: Asian pacific association of gastroenterology and Asian pacific society for digestive endoscopy; APA: Antiplatelet agents; ASA: Acetylsalicylic acid.



**Figure 3 Management of warfarin in elective endoscopic procedures according to international guidelines.** ASGE: American society for gastrointestinal endoscopy; ESGE/BSG: European society of gastrointestinal endoscopy and British society of gastroenterology; APAGE/APSDE: Asian pacific association of gastroenterology and Asian pacific society for digestive endoscopy; INR: International Normalized Ratio; LMWH: Low molecular weight heparin.



**Figure 4 Management of direct oral anticoagulants in elective endoscopic procedures according to international guidelines.** ASGE: American society for gastrointestinal endoscopy; ESGE/BSG: European society of gastrointestinal endoscopy and British society of gastroenterology; APAGE/APSDE: Asian pacific association of gastroenterology and Asian pacific society for digestive endoscopy; DOAC: Direct oral anticoagulant.