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Responsible, Safe, and Effective Use of Biologics in the Management of Low Back Pain: American Society of Interventional Pain Physicians (ASIPP) Guidelines

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Guidelines

Responsible, Safe, and Effective Use of Antithrombotics and Anticoagulants in Patients Undergoing Interventional Techniques: American Society of Interventional Pain Physicians (ASIPP) Guidelines

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From: American Society of Interventional Pain Physicians

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Background: Interventional pain management involves diagnosis and treatment of chronic pain. This specialty utilizes minimally invasive procedures to target therapeutics to the central nervous system and the spinal column. A subset of patients encountered in interventional pain are medicated using anticoagulant or antithrombotic drugs to mitigate thrombosis risk. Since these drugs target the clotting system, bleeding risk is a consideration accompanying interventional procedures. Importantly, discontinuation of anticoagulant or antithrombotic drugs exposes underlying thrombosis risk, which can lead to significant morbidity and mortality especially in those with coronary artery or cerebrovascular disease. This review summarizes the literature and provides guidelines based on best evidence for patients receiving anti-clotting therapy during interventional pain procedures.

Study Design: Best evidence synthesis.

Objective: To provide a current and concise appraisal of the literature regarding an assessment of the bleeding risk during interventional techniques for patients taking anticoagulant and/or antithrombotic medications.

Methods: A review of the available literature published on bleeding risk during interventional pain procedures, practice patterns and perioperative management of anticoagulant and antithrombotic therapy was conducted. Data sources included relevant literature identified through searches of EMBASE and PubMed from 1966 through August 2018 and manual searches of the bibliographies of known primary and review articles.

Results:

- 1. There is good evidence for risk stratification by categorizing multiple interventional techniques into low-risk, moderate-risk, and high-risk. Also, their risk should be upgraded based on other risk factors.
- 2. There is good evidence for the risk of thromboembolic events in patients who interrupt antithrombotic therapy.
- 3. There is good evidence supporting discontinuation of low dose aspirin for high risk and moderate risk procedures for at least 3 days, and there is moderate evidence that these may be continued for low risk or some intermediate risk procedures.

- 4. There is good evidence that discontinuation of anticoagulant therapy with warfarin, heparin, dabigatran (Pradaxa®), argatroban (Acova®), bivalirudin (Angiomax®), lepirudin (Refludan®), desirudin (Iprivask®), hirudin, apixaban (Eliquis®), rivaroxaban (Xarelto®), edoxaban (Savaysa®, Lixiana®), Betrixaban(Bevyxxa®), fondaparinux (Arixtra®) prior to interventional techniques with individual consideration of pharmacokinetics and pharmacodynamics of the drugs and individual risk factors increases safety.
- 5. There is good evidence that diagnosis of epidural hematoma is based on severe pain at the site of the injection, rapid neurological deterioration, and MRI with surgical decompression with progressive neurological dysfunction to avoid neurological sequelae.
- 6. There is good evidence that if thromboembolic risk is high, low molecular weight heparin bridge therapy can be instituted during cessation of the anticoagulant, and the low molecular weight heparin can be discontinued 24 hours before the pain procedure.
- 7. There is fair evidence that the risk of thromboembolic events is higher than that of epidural hematoma formation with the interruption of antiplatelet therapy preceding interventional techniques, though both risks are significant.
- 8. There is fair evidence that multiple variables including anatomic pathology with spinal stenosis and ankylosing spondylitis; high risk procedures and moderate risk procedures combined with anatomic risk factors; bleeding observed during the procedure, and multiple attempts during the procedures increase the risk for bleeding complications and epidural hematoma.
- 9. There is fair evidence that discontinuation of phosphodiesterase inhibitors is optional (dipyridamole [Persantine], cilostazol [Pletal]. However, there is also fair evidence to discontinue Aggrenox [dipyridamole plus aspirin]) 3 days prior to undergoing interventional techniques of moderate and high risk.
- 10. There is fair evidence to make shared decision making between the patient and the treating physicians with the treating physician and to consider all the appropriate risks associated with continuation or discontinuation of antithrombotic or anticoagulant therapy.
- 11. There is fair evidence that if thromboembolic risk is high antithrombotic therapy may be resumed 12 hours after the interventional procedure is performed.
- 12. There is limited evidence that discontinuation of antiplatelet therapy (clopidogrel [Plavix®], ticlopidine [Ticlid®], Ticagrelor [Brilinta®] and prasugrel [Effient®]) avoids complications of significant bleeding and epidural hematomas
- 13. There is very limited evidence supporting the continuation or discontinuation of most NSAIDs, excluding aspirin, for 1 to 2 days and some 4 to 10 days, since these are utilized for pain management without cardiac or cerebral protective effect.

Limitations: The continued paucity of the literature with discordant recommendations.

Conclusion: Based on the survey of current literature, and published clinical guidelines, recommendations for patients presenting with ongoing antithrombotic therapy prior to interventional techniques are variable, and are based on comprehensive analysis of each patient and the risk-benefit analysis of intervention.

Key words: Perioperative bleeding, bleeding risk, practice patterns, anticoagulant therapy, antithrombotic therapy, interventional techniques, safety precautions, pain

Disclaimer: These guidelines are based on the best available evidence and do not constitute inflexible treatment recommendations. Due to the changing body of evidence, this document is not intended to be a "standard of care."

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1.0 Introduction

ith the increasing use of prescription medications to manage thrombosis risk and the ease of availability of over-thecounter and herbal products that mediate or modulate the coagulation cascade (1-21), and techniques to manage chronic pain, interventional pain physicians frequently encounter the challenge of potential risk of bleeding or thrombosis in perioperative management of these patients (1,13,14,20,22-54). Modulation of anticoagulant and antithrombotic therapy during performance of interventional techniques is one of the major clinical decisions often made without precise evidence-based literature to support existing opinions (1,13,14,20,31-54). Leading causes of morbidity and mortality worldwide include cardiovascular and cerebrovascular disease (4,5,55-58). In addition, one of the leading causes of disability and functional impairment across the globe is chronic persistent pain (29,30,59-66). Therefore, overlap between chronic persistent pain and cardiovascular disease has a synergistic impact on physical and psychological health, affecting performance of social responsibilities, including work and family life. Antithrombotic therapy has a clear evidence-based foundation with a favorable risk-benefit profile for prevention and management of cardiovascular disease, including limiting the present and future burden of cardiac or cerebrovascular infarcts (4,5,15-19,55-58,67-81). Of note, a significant proportion of patients with established cerebrovascular, cardiovascular or peripheral vascular disease who are receiving antithrombotic therapy, are commonly in need of interventions including surgery and interventional pain management techniques, despite the debate regarding their safety, clinical and cost effectiveness, and indications with numerous regulations (1,22-30,81-121).

Based on a survey of common practices by a majority of interventional pain physicians, discontinuation of antiplatelet therapy and anticoagulant therapy is common (31). The results of this survey showed discontinuation rate of warfarin therapy 100%, clopidogrel 97%, ticlopidine 96%, Aggrastat or tirofiban 95%, cilostazol 93%, dipyridamole 85%, aspirin 350 mg 60%, aspirin 81 mg 39%, and other nonsteroidal anti-inflammatory drugs (NSAIDs) 39% prior to performing interventional pain management techniques. The majority of physicians accepted an international normalized ratio (INR) of 1.5 or less as a safe level. In another survey performed

in 2012 by the American Society of Regional Anesthesia and Pain Medicine (ASRA), 55% stopped aspirin before spinal cord stimulation trials and implants, and 32% stopped before epidural steroid injections (40). The results of this study showed that only 36% knew that selective serotonin reuptake inhibitors (SSRIs) may lead to a bleeding disorder. Further, 17% utilized different protocols for cervical spine injections as compared with lumbar spine injections. This study (40) also showed that most physicians (88%) expressed the need for pain physicians to communicate with other physicians.

Based on published guidelines, derived by clinical case reports and consensus, a large subset of clinicians report stopping antiplatelet therapy and consider this concept as a standard of care (1,20,33,36-38,122-129). Although the overall incidence of bleeding complications and epidural hematoma in the non-obstetric epidural literature has been reduced, the incidence has been higher with procedures involving cervical and thoracic spine (1,31-42,52-54,129). Of note, the clinical literature reports of incidences of epidural hematoma cases accompanying interventional techniques and neuraxial techniques are increasing rapidly (1,20,31-42,130-149). While some reports indicate a decreasing incidence of bleeding complications related to neuraxial techniques (145), multiple studies have been performed assessing the prevalence and risks related to bleeding complications and epidural hematoma with neuraxial procedures, specifically interventional techniques (1,20,31-42,50,51,138-149).

Significant risks of withdrawing antiplatelet therapy include cardiovascular, cerebrovascular, and peripheral vascular thrombosis. In addition, the risks are higher in chronic pain patients as chronic psychosocial stress causes a hypercoagulable state, as reflected by increased procoagulant molecular fibrinogen (or coagulation factor 7), reduced fibrinolytic capacity and increased platelet activity (40,150-152). Stress has been shown to affect coagulation activity via an influence on the regulation of genes coding for coagulation and fibrinolysis molecules (152) with increase in hormonal levels (153-155) and an underlying surge of catecholamine and cortisol induced hypercoagulability (155). A prior systematic review and meta-analysis explored the hazards of discontinuing or altering aspirin regimens among patients at risk for coronary artery disease (69). Importantly, in patients at moderate to high risk for coronary artery disease, withdrawal or noncompliance

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with aspirin therapy was associated with 3-fold higher risk of major adverse cardiac events and the risk was magnified in patients with coronary stents. The findings support the recommendation that aspirin discontinuation in this patient population should be advocated only under circumstances where the risk of adverse outcomes caused by bleeding risk clearly outweighs that of catastrophic atherothrombotic events. These findings have been confirmed in later studies (70-78,150). Conversely, recently published large-scale evidence (15-19) shows lack of benefit of aspirin for primary prevention of cardiovascular events, but aspirin therapy is associated with increased bleeding episodes. Thus, current evidence suggests that the risks of coronary events related to patients abstaining from their antiplatelet medications during the perioperative period are more serious when compared to the risks of continuing antiplatelet therapy through the perioperative period.

Multiple publications (1,32,33,42,46,138,140) have also supported the concept of continuing antiplatelet/ anticoagulant agents in patients undergoing various interventional pain procedures in light of identical complication rates as compared to the patients who stop taking these for a particular recommended period. It should be noted that there are only a few clinical reports of an epidural hematoma available in patients undergoing interventional therapies for chronic pain, which included patients both continuing and discontinuing antithrombotic therapy.

The American Society of Interventional Pain Physicians (ASIPP) has developed multiple guidelines, both for interventional techniques and opioids (29,66,82). Consequently, the present investigation has been undertaken to develop responsible, safe, and effective guidelines for patients on antithrombotic and anticoagulant therapy undergoing interventional techniques. These guidelines offer an overview of the current literature applicable to antithrombotic and anticoagulant therapy and the related complications, with incorporation of various aspects of the current national and international recommendations.

2.0 Methods

2.1 Rationale

Interventional pain management physicians provide a significant number of multidisciplinary strategies including physical therapy, behavioral treatments, pharmacological, and interventional management to patients with chronic pain. The National Uniform

Claims Committee (NUCC) defined interventional pain management as the discipline of medicine devoted to the diagnosis and treatment of pain-related disorders, principally with the application of interventional techniques in managing subacute, chronic, persistent, and intractable pain, independently or in conjunction with other modalities of treatment (156). In addition, the Medicare Payment Advisory Commission (MedPAC) defined interventional pain management techniques as minimally invasive procedures including, percutaneous precision needle placement with placement of drugs in targeted areas or ablation of targeted nerves; and some surgical techniques such as laser or endoscopic discectomy, intrathecal infusion pumps and spinal cord stimulators, for the diagnosis and management of chronic, persistent or intractable pain (157).

Chronic spinal pain is a complex and multifactorial phenomenon. Interventional pain physicians are familiar with various pathophysiologic, anatomic, and technical aspects of multiple interventional techniques applied in managing chronic pain.

2.2 Objectives

The objectives of these guidelines are to synthesize the available evidence on the safety, as well as adverse effects of antithrombotic and anticoagulant therapy for patients undergoing interventional techniques in the treatment of chronic spinal pain, and provide a rational and systematic approach to their application.

2.3 Adherence to Trustworthy Standards

In preparation of these guidelines to develop responsible, safe, and effective guidelines for patients on antithrombotic and anticoagulant therapy undergoing interventional techniques, the Institute of Medicine (IOM) standards and the National Guideline Clearinghouse Extent Adherence to Trustworthy Standards (NEATS) instruments were followed (158-160). The NEATS instrument was developed and tested as a tool to be used by the trained staff at the Agency for Healthcare Research and Quality (AHRQ) National Guideline Clearinghouse to provide assessment focused on adherence.

2.3.1 Disclosure of Guideline Funding Source

Responsible, safe, and effective guidelines for patients on antithrombotic and anticoagulant therapy undergoing interventional techniques were commissioned, prepared, edited, and endorsed by ASIPP without external funding sought or obtained.

Table 1. Qualitative modified approach to grading of evidence.

Level I	Strong	Evidence obtained from multiple relevant high quality randomized controlled trials for effectiveness
Level II	Moderate	Evidence obtained from at least one relevant high quality randomized controlled trial or multiple relevant moderate or low quality randomized controlled trials
Level III	Fair	Evidence obtained from at least one relevant high quality nonrandomized trial or observational study with multiple moderate or low quality observational studies
Level IV	Limited	Evidence obtained from multiple moderate or low quality relevant observational studies
Level V	Consensus based	Opinion or consensus of large group of clinicians and/or scientists for effectiveness as well as to assess preventive measures, adverse consequences, effectiveness of other measures.

Modified from: Manchikanti et al. A modified approach to grading of evidence. Pain Physician 2014; 17:E319-E325 (161).

2.3.2 Disclosure and Management of Financial Conflicts of Interests

Potential conflicts of interest for all panel members within the last 5 years were evaluated prior to finalizing these guidelines. However, the panel members with potential conflicts were instructed by the panel and recused from related discussion or preparation of the guidelines and these members agreed not to discuss any aspect of the guideline with the industry before data publication. Further, conflicts of interests were included based on interest confluence extending beyond financial relationships including personal experience, practice patterns, academic interests and promotions.

2.3.3 Composition of the Guideline Development Group

A panel of experts, convened by ASIPP, reviewed the evidence and formulated recommendations for interventional techniques in patients receiving antiplatelet and anticoagulant therapy. The panel was instructed to assess the evidence pertaining to all important aspects of therapy and related complications.

The panel provided a broad representation of academic and non-academic clinical practitioners with interest in interventional techniques in patients receiving antiplatelet and anticoagulant therapy.

The panel composition was multidisciplinary including methodologists (e.g., epidemiologists, statisticians, and health services researchers) with experience in research and the conduct of systematic reviews. These members were voting members of the guideline development groups.

2.4 Evidence Review

Responsible, safe, and effective guidelines for patients on antithrombotic and anticoagulant therapy undergoing interventional techniques were developed utilizing evidence review, incorporating guidelines by other organizations and agencies, if available, and developing consensus among the panel members.

The current guidelines offer recommendations based on scientific evidence and informed expert opinion.

2.4.1 Grading or Rating the Quality or Strength of Evidence

This grading of evidence is based on randomized controlled trials (RCTs), observational studies, and other clinical reports based on availability. In addition, systematic reviews, meta-analyses, and evidence developed by other guidance are also given high importance with critical analysis. The recommendations have been developed using principles of best evidence synthesis developed by the Cochrane Review, incorporating multiple guidelines modified by ASIPP as shown in Table 1 (161).

This methodology specifies a level of scientific evidence and offers a transparent approach to grading quality of evidence and strength of recommendations. AHRQ also has recommended a similar strength of recommendation as shown in Table 2 (159,160).

2.4.2 Assessment and Recommendations of Benefits and Harms

These guidelines intend to clearly describe the potential benefits and harms and explicitly link the information to specific recommendations.

2.4.3 Evidence Summary of Recommendations

Guideline supporting documents summarize the relevant supporting evidence and explicitly link this information to recommendations.

2.4.4 Rating or Grading the Strength of Recommendations

IOM standards demand that for each recommenda-

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Table 2. Guide for strength of recommendations.

Rating for Strength	of Recommendation	
Strong	There is high confidence that the recommendation reflects best practice. This is based on: a) strong evidence for a true net effect (e.g., benefits exceed harms); b) consistent results, with no or minor exceptions; c) minor or no concerns about study quality; and/or d) the extent the panelists' agreement. Other compelling considerations (discussed in the guideline's literature review and analyses) may also warrant a strong recommendation.	
Moderate	There is moderate confidence that the recommendation reflects best practice. This is based on: a) good evidence for a true net effect (e.g. benefits exceed harms); b) consistent results, with minor and/or few exceptions; c) minor and/or few concerns about study quality; and/or d) the extent of panelists' agreement. Other compelling considerations (discussed in the guideline's literature review and analyses) may also warrant a moderate recommendation.	
Weak	There is some confidence that the recommendation offers the best current guidance for practice. This is based on: a) limited evidence for a true net effect (e.g., benefits exceed harms); b) consistent results, but with important exceptions; c) concerns about study quality; and/or d) the extent of panelists' agreement. Other considerations (discussed in the guideline's literature review and analyses) may also warrant a weak recommendation.	

Source: National Guideline Clearinghouse Extent Adherence to Trustworthy Standards (NEATS) instrument (159).

tion, a rating of the strength of the recommendation related to benefits and harms, available evidence, and the confidence in the underlying evidence should be provided (158).

2.4.5 Specificity of Recommendations

Guideline recommendations in this manuscript may not be very specific and occasionally may be ambiguous, on what actions should or should not be taken in various situations of antithrombotic and anticoagulant therapy during performance of interventional techniques due to the lack of extensive literature and discordant opinions.

2.5 Evidence Synthesis and Analysis

This comprehensive review was performed with appropriate literature review and assessment. Due to the complexity of the subject and lack of systematic assessment of bleeding complications and epidural hematomas in the literature, a systematic review and meta-analysis of RCTs and/or observational studies may not be feasible (158,162-164). Consequently, these guidelines focus on the available literature in reference to epidemiologic studies, observational reports, and case reports. Further, appropriate methods of data extraction and methodologic quality assessment will be applied.

2.5.1 Data Sources

The literature search was strategized based on multiple available sources including PubMed, the Cochrane library, Google searches, the U.S. Food and Drug Administration (FDA), the Centers for Disease Control and Prevention (CDC), and controlled clinical trials. The search was conducted over a time frame spanning from 1966 to August 2018.

2.5.2 Search Strategy

The search strategy focused on antithrombotics, anticoagulants, antiplatelet agents, thrombotic events, cardiovascular events, cerebrovascular events, interventional techniques, bleeding complications, and epidural hematoma following interventional techniques.

2.5.3 Study Selection

The review focused on all types of reports and includes case reports and reviews.

- Abstracts were screened in an unblinded standardized manner by 2 review authors for all identified studies.
- Full text for all articles with relevance to the topic was retrieved for comprehensive review.

2.5.4 Data Extraction and Methodological Quality Assessment

Data extraction and methodologic quality assessment was performed by at least two of the review authors independently, in an unblinded standardized manner. The quality assessment of each individual article used in this analysis was performed by utilizing the Cochrane review criteria (Appendix Table 1) (165), Interventional Pain Management techniques Quality Appraisal of Reliability and Risk of Bias Assessment (IPM-QRB) criteria (Appendix Table 2) (166) for RCTs, and Interventional Pain Management Techniques - Quality Appraisal of Reliability and Risk of Bias Assessment for Nonrandomized Studies (IPM-QRBNR) (Appendix Table 3) (167). Utilizing the Cochrane review criteria (165) or IPM-QRB criteria (166), or IPM-QRBNR (167), studies meeting the inclusion criteria with a score of at least 9 to 13 for the Cochrane review criteria or 32 to 48 for the IPM-QRB and IPM-QRBNR criteria were considered high quality; while 5 to

8 or 16 to 31 were considered moderate quality; whereas those with scores of less than 5 or 16 were considered low quality. Due to the scarcity of the literature, all studies were included in data synthesis or analysis.

All systematic reviews, non-systematic reviews, all types of studies, and case reports were reviewed by 2 authors. Any disagreements between the reviewers were resolved by a third author and consensus. If there were any conflicts of interest with a manuscript (i.e., authorship) the review authors were recused from assessment and analysis.

2.5.5 Analysis of Evidence

The analysis of the evidence was performed based on best-evidence synthesis and was modified and collated using multiple available criteria, including the Cochrane Review criteria and the United States Preventive Task Force (USPSTF) criteria as illustrated in Table 1 (161). The analysis was conducted using 5 levels of evidence ranging from strong to opinion- or consensus-based. The results of best evidence as per grading were utilized. At least 2 of the review authors independently, in an unblinded, standardized manner, analyzed the evidence. Any disagreements between reviewers were resolved by a third author and consensus was attained. If there were any conflicts of interest (e.g., authorship), the reviewers of interest were recused from assessment and analysis.

2.5.6 Meta-analysis

The meta-analysis was planned to be performed using the Comprehensive Meta-analysis version 3.0 (Biostat Inc., Englewood, NJ). Heterogeneity will be interpreted through I² statistic.

Random-effects model (single-arm) meta-analysis was planned to assess net changes in the same outcome variable (168-170). Heterogeneity among the effect sizes of individual studies will be assessed using the I² index and Q statistic. Heterogeneity analyzed with the I² statistic was defined as low (25%–50%), moderate (50%–75%), or high (>75%).

All analyses were based on if antithrombotics or anticoagulants were utilized or stopped. Meta-analysis was performed only when at least 3 studies were available and included an appropriate sample size of at least 10 for nonrandomized studies.

2.6 External Review

Guidelines have been subjected to external peer review as per the policies of the publishing journal, *Pain Physician*.

2.7 Updating Guidelines

The responsible, safe, and effective guidelines for patients on antithrombotic and anticoagulant therapy undergoing interventional techniques will be updated in a window of 5 years, based on significant changes in the evidence, public policy, or adverse events before January 2024.

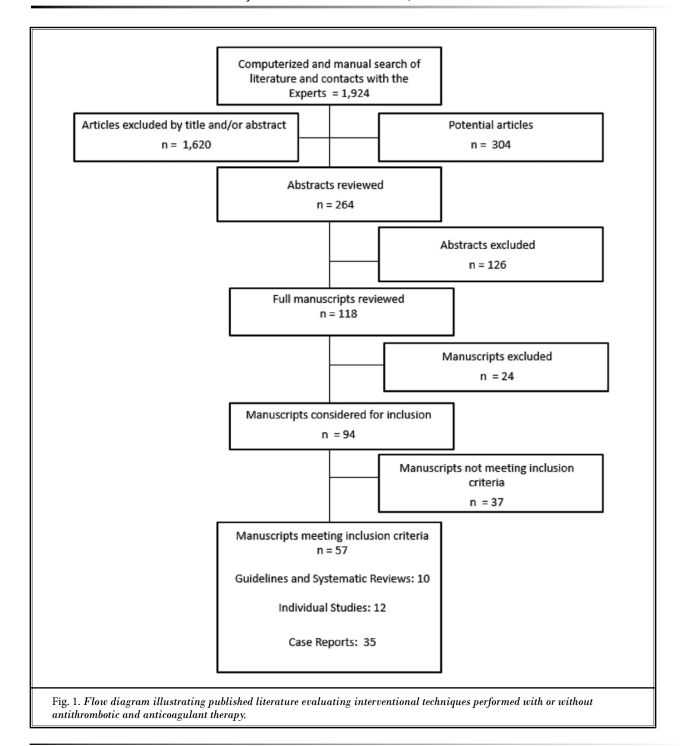
3.0 RESULTS

Our search strategy utilizing the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) flow diagram as shown in Fig. 1, illustrated published literature evaluating interventional techniques performed with or without antithrombotic and anticoagulant therapy. We identified multiple systematic reviews and guidelines (1,20,28,36,37,39,40,82,171-174) and multiple individual manuscripts which met the criteria for inclusion (175-212). Due to very few studies with persistent high variability, conventional meta-analysis or single-arm meta-analysis was feasible.

3.1 Reviews and Guidelines

Among the 12 manuscripts describing guidelines and reviews, 2 manuscripts were related to ASIPP guidelines (1,82). The present guidelines are based on these guidelines published in 2013. Other guidelines of high importance are of the ones published by the ASRA (40). These guidelines have provided extensive descriptions with comprehensive review of the literature; however, these have been considered as more aggressive about stopping the antithrombotics than other available literature (36). Nevertheless, these are commonly utilized guidelines in the United States. Multiple other manuscripts include the guidelines from across the globe and reviews. A review by Oprea et al (20) described risk stratification, perioperative and periprocedural management of the patient receiving anticoagulant therapy.

In addition, there were also guidelines related to the Neurostimulation Appropriateness Consensus Committee with recommendations on bleeding and coagulation management in neurostimulation devices (39). The guidelines from ASRA (40) and guidelines for neuromodulation procedures (39) stratified procedures into various risk categories. The guidelines by ASRA were revised which was prompted by reports of percutaneous lead placement with development of epidural hematoma in patients on aspirin. Prior to these recommendations, low dose aspirin was considered as safe and was not stopped during interventional techniques including spinal cord stimulation.



Breivik et al (36) provided a comprehensive review of reducing the risk of spinal hematoma from spinal and epidural pain procedures. They provided recommendations for interventional techniques with updated guidance.

In addition, in a quasi-systematic review assessing the risks and benefits of ceasing or continuing anticoagulant medication for image-guided procedures for spine pain (33), the authors evaluated 14 manuscripts which provided applicable evidence. The results showed procedures involving interlaminar access carried a nonzero risk of hemorrhagic complications, regardless of whether anticoagulants were ceased or continued. For other procedures, hemorrhagic complications have not been reported, and case series indicated that they are safe when performed in patients who continue anticoagulants. The results of this review identified reports of adverse effects of ceasing anticoagulants, with serious consequences, including death. The authors of this manuscript concluded that other than for interlaminar procedures, the evidence does not support the view that anticoagulant and antiplatelet medication must be ceased before image-guided spine pain procedures.

3.2 Prevalence and Risk Assessment of Studies

Multiple studies have been published assessing the prevalence, as well as the risk of bleeding with interventional techniques (31-34,37,41,42,50-54).

In a survey of practice patterns among interventional pain physicians in 2012, Manchikanti et al (31) showed that the majority of physicians discontinued antithrombotic agents; however, this study also showed that there were a significantly higher number of complications related to thromboembolic events of a total of 162 compared to hemorrhagic complications of a total of 55 in this population.

In a prospective evaluation of bleeding risks for interventional techniques in chronic pain, Manchikanti et al (32) assessed the rates of adverse events in patients undergoing interventional techniques on antithrombotic therapy with cessation or without cessation and compared them to a group of patients without antithrombotic therapy. While the results showed differences in milder complications, there were no reports of hemorrhagic complications requiring any type of treatment. In this assessment, the authors studied all types of procedures with 1,227 of 1,831 continuing aspirin compared to 604 of 1,831 discontinuing them. Similarly, they also studied 100 patients on clopidogrel with continuation, whereas, 226 patients were discontinued. Further, there were 128 patients with aspirin and other agents with continuation and 151 were discontinued. The procedures performed included cervical epidural injections with continued aspirin in 249 patients, thoracic epidural in 30 patients, lumbar interlaminar epidural in 128 patients, lumbar transforaminal in 144 patients, whereas 528 patients for caudal epidural injections, and 148 for percutaneous adhesiolysis. In reference to clopidogrel, it was continued in 10 patients undergoing cervical epidural, one patient with thoracic

epidural, 14 patients with lumbar epidural, 44 patients with caudal epidurals, 10 with lumbar transforaminal epidural, and 21 with percutaneous adhesiolysis. There were a large number of facet joint interventions and other treatments.

van Helmond et al (51), in a retrospective review, assessed the safety of low to intermediate risk spine procedures in patients with continued antithrombotic therapy. They identified 490 patients of a total of 2,204 patients on antithrombotic medications which included aspirin (N=275), P2Y12 inhibitors (N = 129), warfarin (N = 62), heparin (N = 10), factor Xa inhibitors (N = 55), and dipyridamole (N = 1). The procedures included facet joint nerve blocks and facet joint radiofrequency in all 3 regions and sacroiliac joint injections. The authors concluded that there were no hemorrhagic complications in performing these procedures.

Goodman et al (50) studied the role of anticoagulant and antiplatelet management for spinal procedures in a prospective descriptive study. Out of 4,253 procedures, 197 of them were performed in 74 patients on antiplatelet/anticoagulants, they reported no clinically evident bleeding events for lumbar transforaminal epidural injections (N=90), facet joint injections (N=62), lumbar intradiscal procedures (N=11), lumbar sympathetic blocks (N=3), sacroiliac injections (N=5), or in 26 radiofrequency neurotomy procedures. However, similar to van Helmond et al (51), Goodman et al (50) have not performed any epidural injections on patients with continuation of antiplatelet therapy.

Endres et al (42) assessed risks of continuing or discontinuing anticoagulants for patients undergoing interventional procedures. They reported no complications attributable to anticoagulants in 4,766 procedures in which anticoagulants were continued. They mainly performed lumbar transforaminal injections, lumbar medial branch blocks, trigger point injections, and sacroiliac joint blocks, and they concluded that continuation of anticoagulant therapy seems to be safe. They performed interlaminar epidural injections in 25 patients with continuation of Warfarin and 15 patients with continuation of clopidogrel. In addition, they reported 171 patients undergoing sacroiliac joint blocks on Warfarin, 227 patients undergoing trigger point injections, 40 patients with trochanteric bursa injections, and 87 patients with hip joint injections, they also reported 81 patients with clopidogrel undergoing sacroiliac joint blocks, 214 patients with trigger point injections, 50 patients with trochanteric bursa injections, and 52 patients with hip joint injections.

Warner et al (54) in a manuscript describing bleeding and neurological complications in 58,000 interventional pain procedures showed that preprocedural aspirin or nonsteroidal antiinflammatory drug therapy was prevalent in 17,825 procedures or 30.7% of the procedures without significant bleeding complications. Out of total 58,066 procedure performed in the study; 22.4% of the procedures were performed with perioperative administration of aspirin within 7 days, 12.1% of the patients with administration of nonsteroidal anti-inflammatory drugs within 7 days, 1.6% of clopidogrel within 7 days. The study also included 3,880 lumbar epidural injections, 304 thoracic interlaminar injections with a large number of epidural injections with over 50% of the epidural injections not assigned to a region. They concluded that bleeding complications were rare in patients undergoing low or intermediate risk pain procedures even in the presence of antiplatelet medication.

Lagerkranser (52) and Lagerkranser and Lindquist (53) have published extensive review of neuraxial blocks and spinal hematoma in 2 parts from 1994 to 2015 covering demographics, risk factors, diagnosis, treatment, and outcome. They also considered previous reviews published in 1992 (175), 1994 (176), and 1996 (177) case reviews, analyzing 29, 61, and 51 cases of spinal hematoma after neuraxial blocks, respectively, between 1906 and 1996. They identified multiple changes in the perioperative management of surgical patients, thromboprophylaxis with multiple newer agents, and multiple published guidelines. They identified 147 published articles, case reports or letters, describing altogether 166 cases relevant for the review. They showed that 64 cases were from Europe, 53 from North America, 4 from South America, 39 from Asia and 6 from Australia. They reported hematoma formation secondary to multiple types of neuraxial blocks, including spinal injections, epidural injections, combined spinal and epidural injections, and spinal cord stimulators. They reported 49 lumbar injections with 12 single shots, 46 thoracic injections all with continuous and 6 cervical locations with 5 single shots secondary to epidural injections. A large number of hematomas appeared after removal of the catheter. In managing chronic pain with epidural injections, they identified 21 hematomas, 17 (5 cervical, 4 thoracic, and 8 lumbar) after epidural injections, and 4 after percutaneous application of spinal cord stimulators. However, they have not identified the number of patients developing hematoma with appropriate cessation of antithrombotic therapy based on recommended

guidelines. Overall, they showed that 37% of the patients who were not on antihemostatic drugs, whereas 63% were on antihemostatic drugs with 47 of the cases, receiving more than one antihemostatic drug, and 12 receiving 3 such drugs. Further, they also had 6 reports which were indeterminate. Consequently, the number of patients without antithrombotic therapy, but with hematoma formation seems to be almost 40%.

Lagerkranser (52) showed that an annual average of 7.5 published cases of spinal hematoma in the years 1994 to 2015, compared to an average of only 2.5 case reports per year from 1976 to 1993 (176). They also identified that there has been a transition from a male to female dominance among patients suffering from post-neuraxial blockade and spinal hematoma particularly among the elderly women. In addition, guidelines issued around the turn of the century have not been followed by a reduced number of published case reports, even though adherence to the guidelines has increased during the last decade.

The authors identified spinal stenosis as the most common spinal disease in 14 cases with spinal disease reported in 37 cases of 166 cases (52). They identified bloody tap at the introduction of a neuraxial needle or catheter as a major risk factor, but multiple attempts to reach the spinal canal do not seem to increase the risk of spinal hematoma. Their results also showed that 80% of the patients developing spinal hematoma had severe neurological symptoms with paresis or paralysis. When compared over time, outcomes have improved significantly (52). The results showed that among patients subjected to surgical evacuation of the hematoma, outcomes were best if surgery was performed within 12 hours from the first sign of motor dysfunction. However, even patients operated after more than 24 hours had relatively favorable outcomes. Further, outcomes after surgical evacuation of the epidural hematoma were satisfactory, compared to subdural hematoma, which had poor outcomes. They recommended that suspicion of spinal hematoma calls for the consultation of surgeon without delay. Magnetic resonance imaging (MRI) was the recommended diagnostic tool. Surgical evacuation within 12 hours from the sign of motor dysfunction seems to lead to the best outcome, even though many patients operated as late as after more than 24 hours did regain full motor function (52).

Moeschler et al (34) assessed bleeding complications in patients undergoing percutaneous spinal cord stimulator trials and implantations with retrospective review from 2005 through 2014 of all patients, with assessment of 642 percutaneous spinal cord procedures, 346 spinal cord stimulator trials, 255 spinal cord stimulation implantations, and 41 revision surgeries. Patients had received aspirin or NSAIDs within 7 days of needle placement for 101 procedures (15.7%). There were no bleeding or neurological complications identified in this cohort. Consequently, this study showed that there were no risks associated with patients receiving aspirin or other NSAIDs within 7 days of needle placement in approximately 16% or 100 procedures; however, they warned that further investigations with larger numbers are needed to better define the relationship between periprocedural aspirin and NSAID utilization and bleeding complications.

Petraglia et al (41) also studied the incidence of spinal cord injury in implantation of percutaneous and paddle electrodes for spinal cord stimulation, with analysis of nationwide data, with inclusion of 8,326 patients. They showed overall incidence of spinal cord injury of 2.1%, with incidence of spinal hematoma of 0.71%, which was seen in 59 patients with 41 in percutaneous group with 0.75% and 18 in paddle lead group with 0.63%. They concluded that while reversible complications may be more common, spinal cord injury as a result of spinal cord stimulation is uncommon. Even though this was a large database, they recommended for more studies to further characterize the mechanisms of injury and outcomes in these patients. They also assessed odds ratio for using antiplatelet or anticoagulant medications within 30 days prior to spinal cord stimulation procedures. They found that the odds of spinal cord injury increased by 2.4. If the patient took any type of anticoagulant medication within 30 days prior to the procedure they found no significant increase in the odds of spinal cord injury for patients taking aspirin, clopidogrel, or enoxaparin alone.

LaVallee et al (35) studied the prevalence of bleeding complications following ultrasound-guided botulinum toxin injections in patients on anticoagulation or antiplatelet therapy. They reviewed 328 ultrasound guided intramuscular procedures performed in 15 patients with the predominant indication for chemodenervation being spastic paresis secondary to stroke. Of 328 procedures, only 2 subclinical hematomas were detected, resulting in a bleeding complication rate of 0.61% in this patient population.

Table 3 describes the studies assessing the risk of thrombosis and bleeding with interventional pain management techniques. All but 2 studies in this assessment are related to performing interventional tech-

niques without cessation of antithrombotic therapy (30-34,37,39,41,42,50,51). Only 2 studies (31,52) were related to an online physician survey and study of spinal hematoma with neuraxial blocks. Among all the studies, only 2 studies included epidural injections (32,42). All others have performed a large proportion of procedures with low risk or intermediate risk including transforaminal epidural injections. As shown in Table 4, interlaminar procedures were performed to a large extent by Manchikanti et al (32) with inclusion of patients undergoing cervical epidural, thoracic epidural, lumbar epidural, caudal percutaneous adhesiolysis with aspirin, warfarin, clopidogrel, and aspirin plus other drugs. Endres et al (42) also studied lumbar interlaminar epidural injections in 25 patients with continuation of warfarin and 15 patients with continuation of clopidogrel and 2 patients with continuation of cilostazol and enoxaparin. Lagerkranser (52) also reported 37% of the patients who were not on antithrombotics.

Based on the evidence presented from these studies, it appears that there is no significant difference whether antithrombotic therapy is discontinued or continued in reference to the bleeding. A majority of the authors have studied intermediate and low risk procedures without the inclusion of epidural injections. The only one study reviewing spinal cord stimulation also showed lack of increased risk with continuation of aspirin and other NSAIDs.

3.3 Reports of Thromboembolic Events

There have been multiple reports of thromboembolic events with discontinuation of antithrombotics and anticoagulants prior to performance of interventional procedures. Manchikanti et al (31) in assessment of practice patterns of perioperative management of antiplatelet and anticoagulant therapy in interventional pain management reported 162 thromboembolic events compared to 55 serious bleeding complications from epidural hematomas. This study showed thromboembolic events were 3 times more frequent than bleeding complications. Further, they also showed bleeding complications from epidural hematomas were similar whether antiplatelet therapy was continued or discontinued with an occurrence of 26 versus 29 respectively; in this survey the sample sizes were not provided. Consequently, it is difficult to assess exact risk of bleeding complications and similarly thromboembolic events.

Endres et al (42) reported 9 patients with thromboembolic events after cessation of anticoagulant therapy of 1,626 procedures. These complications included 2

Specifical Studies assessing the risk of thrombosis and bleeding with interventional pain management techniques.

	Author Conclusions of the Review	This study essentially shows that even though there is no evidence of increased risk of epidural hematoma, the majority of physicians discontinue antiplatelet therapy despite increased risk of thromboembolic complications.	This study essentially showed that there is no significant difference in bleeding patterns whether antithrombotic agents are continued or discontinued except for warfarin with no fatal incidents in a large proportion of patients.	Even though authors have shown no complications with continuation of NSAIDs or aspirin within 7 days in approximately 16% of the patients, the study consists of a small number of patients. Further, this agrees with older guidance of continuation of NSAIDs and aspirin in the perioperative period, even though it is contradictory to more recent guidelines (40).	This is a large database in the United States; however, the study was up from 2000 to 2009. There seems to be exponential increase of spinal cord stimulator placements since 2009; consequently, this data may not reflect present literature.
Sement techniques.	Conclusions by the Study Authors	The authors concluded that clinicians must balance the risks of thromboembolism and bleeding in each patient prior to the routine discontinuation of antiplatelet therapy.	Of the 1,831 patients receiving aspirin 604 discontinued and 1,227 continued and all of them received epidural injections including cervical, thoracic, lumbar interlaminar, and caudal epidural injections. Of the total of 3,251 patients undergoing facet joint interventions 1,084 discontinued and 2,167 continued. Of the total 326 patients undergoing epidural injections on clopidogrel 226 discontinued and 100 continued with patients undergoing all types of epidural injections including cervical, thoracic, and lumbar interlaminar epidural injections. However, for cervical epidural injections a large proportion discontinued (67) versus continued (10). For facet joint interventions, of the 601 patients of oppidegrel 369 patients discontinued and 232 patients continued. There was no clinical or statistical difference in any of the major aspects of bleeding.	Although the incidence of epidural hematoma is low, the development of bleeding complications following SCS lead placement can be devastating. In the present investigation, we identified no cases of epidural hematoma following percutaneous SCS lead placement, including more than 100 patients receiving aspirin or NSAIDs. Future investigations with larger numbers are needed to better define the relationships between periprocedural aspirin and NSAID utilization and bleeding complications.	Authors concluded that this study showed overall a low incidence supporting that SCS is a safe procedure.
able 5. Statutes assessing are ton of an one observing wan take behavioral pain management techniques	Results	The results illustrated an overwhelming pattern of discontinuing antiplatelet and warfarin therapy as well as aspirin and other NSAIDs prior to performing interventional pain management techniques. However, thromboembolism complications were 3 times more prevalent than epidural hematomas (162 versus 55 events).	In this study, one-quarter (3,087) of patient encounters undergoing interventional pain management procedures, were on antithrombotic therapy. Antithrombotic therapy was continued in 55% of the patients or 1,711 encounters, whereas, it was discontinued in 45% of the patients or 1,376 in 45% of the patients or 1,376 in 45% of the patients or 1,376 sercounters. There was no difference in significant side effect rate with or without continuation of antithrombotic therapy:	They performed 642 percutaneous spinal cord stimulation procedures, trial, revision, or implantation in 421 patients. No major bleeding complications. In patients who had taken aspirin or patients who had taken aspirin or NSAIDS within 7 days of procedure. There were no bleeding or neurological complications identified in this cohort.	Overall incidence of spinal hematoma was 59 or 0.71% with percutaneous contributing to 41 or 0.75% incidence versus paddle lead contributing to 18 or 0.63% incidence. There was no significant difference between the groups with patients in the percutaneous group and in the paddle group experiencing a spinal hematoma.
o man cacocamo na fo usa can Sancescen e	Methods	An online physician survey of antithrombotics with complications with or without discontinuation of various antithrombotics and anticoagulants.	The prospective evaluation of measurable outcomes of intravascular entry of the needle, bruising, local bleeding, profuse bleeding, local hematoma, oozing, and postoperative soreness.	A total of 642 percutaneous SCS procedures were performed on 421 unique patients, including 346 SCS trials, 255 SCS implantations, and 41 revision surgeries. Patients had received aspirin or NSAIDs within 7 days of needle placement for 101 procedures (15.7%).	Authors conducted a retrospective review of the Thomson Reuters MarketScan Database of all patients that underwent percutaneous or paddle SCS implantation from 2000 to 2009, to assess the incidence of spinal cord injury including spinal hematoma. Of the 8,256 patients meeting inclusion criteria for the study, 5,458 were percutaneous and 2,868 were paddle leads. The overall incidence of spinal cord injury was 177 or 2,13% with percutaneous lead placement attributing to 128 or 2,35% incidences versus paddle leads contributing to 49 or 1,71% incidence.
Taore 7. Stage	Study/Year	Manchikanti et al, 2012 (31)	Manchikanti et al, 2011 (32)	Moeschler et al 2016 (34)	Petraglia et al, 2016 (41)

Table 3 (cont.). Studies assessing the risk of thrombosis and bleeding with interventional pain management techniques.

Study/Year	Methods	Results	Conclusions by the Study Authors	Author Conclusions of the Review
Endres et al, 2017 (42)	The study was performed as an observational study in a private practice in which some partners continued anticoagulants while other partners routinely discontinued anticoagulants. They studied 4,766 procedures in which anticoagulants were continued and 2,296 procedures in which anticoagulants were discontinued.	No complications attributable to anticoagulants were encountered in 4,766 procedures in which anticoagulants were continued; however, in 2,296 procedures in which anticoagulants were discontinued, according to the guidelines, 9 patients suffered serious morbidity, including 2 deaths.	Lumbar transforaminal epidural injections, lumbar facet joint nerve blocks, trigger point injections, and sacroiliac joint blocks appear to be safe in patients who continue anticoagulants. In patients discontinuing anticoagulants, serious complications are observed in a low proportion of patients with 0.2%.	This is an observational study in a large number of patients, however, interlaminar epidural injections or other high risk procedures were not included. Further, authors conclusions are appropriate correlating with other conclusions that risk of serious complications of discontinuing anticoagulant therapy, even though this risk is low at 0.2%.
Warner et al, 2017 (54)	Retrospective cohort of adult patients undergoing low and intermediate risk pain procedures were assessed from 2005 through 2014 at a single academic retriary care center. A total of 58,006 procedures were performed on 24,590 patients. Antiplatelet therapy included preprocedural aspirin or nonsteroidal antiinflammatory drug therapy in 17,825 procedures comprising of 30.7%.	Of the 58,066 procedures, 16 procedures were associated with perioperative red blood cell transfusion of 0.03%, with no difference based on preprocedural nonsteroidal antiinflammatory drug, including aspirin, or other anticoagulation use. The study results showed that 5 patients (0.009%) had neurologic complications requiring further evaluation. Of the 5 patients, 2 were likely realted to procedural bleeding. 22% of the patients received aspirin within 7 days, 12% of the patients received nonsteroidal antiinflammatory drugs within 7 days, 20 colpidogrel within 7 days and Coumadin within 7 days in 3% of the patients. They also maintained INR of 1.0. The study also included 3,880 lumbar epidural injections with a large numbar of epidural injections with a large number of epidural injections with voer 50% of the epidural injections not assigned to a region.	Authors concluded that bleeding complications are rare in patients undergoing low or intermediate pain procedures, even in the presence of antiplatelet medications.	Authors studied low and intermediate risk procedures in patients on nonsteroidal antiinflammatory agents including aspirin. Of these, 22% of the patients received aspirin within 7 days. The study also included 3.880 lumbar epidural injections, 304 thoracic interlaminar injections with a large number of epidural injections with over 50% of the epidural injections not assigned to a region.
Goodman et al, 2017 (50)	A prospective descriptive evaluation of patients undergoing interventional pain procedures on various antiplatelet and anticoagulant agents at a single interventional physiatry practice.	Overall incidence of spinal epidural hematoma for all procedures studied was one in 4,047 procedures (0.02%, 95% CI ½ 0.000-0.15%). No thromboembolic events (myocardial infarction, cerebrovascular accident, or critical limb ischemia) were observed within 24–48 hours after spinal injection for all patients in the study.	Continuation of clopidogrel or warfarin for lumbar transforaminal epidural and posterior-approach facet joint injections may be reasonable. Interlaminar injections carry greater bleeding risk and merit consideration of holding anticoagulant/antiplatelet agents.	Authors have not provided any conclusive evidence if one procedure is safer to perform than the other while patient continues the anti-thrombotic agents.

Table 3 (cont.). Studies assessing the risk of thrombosis and bleeding with interventional pain management techniques.

Study/Year	Methods	Results	Conclusions by the Study Authors	Author Conclusions of the Review
van Helmond et al, 2017 (51)	Retrospective review of the safety of low to intermediate risk spine procedures in patients with continued antithrombotic therapy. In this study authors identified 490 patients of total of 2,204 patients on antithrombotic medications which included aspirin, P2Y12 inhibitors, warfarin, heparin, factor Xa inhibitors, and dipyridamole. The procedures and dipyridamole. The procedures included facet joint nerve blocks and facet joint radiofrequency in all spine regions and sacroiliac joint injections.	There were no hemorrhagic complications in performing these procedures.	Authors concluded that there were no hemorrhagic complications in performing these procedures and they were safe.	The results are in a small number of patients with low risk and intermediate risk procedures of facet joint interventions and sacrolliac joint interventions without inclusion of high risk procedures of interlaminar epidural injections.
Horlocker et al, 2002 (37)	A prospective study of 1,214 epidural steroid injections in ambulatory pain centers. 32% of the patients were receiving nonsteroidal antiinflammatory agents including 34 patients on multiple medications. Aspirin was the most common NSAID and was noted by 158 patients including 140 patients on 325 mg or less per day.	There were no major hemorrhage complications with spinal hematomas. There were minor hemorrhagic complications with blood during the needle or catheter placement in 5% of the patients. NSAIDS did not increase the frequency of minor hemorrhagic complications. Increased age, needle gauge, needle approach, needle insertion at multiple interspaces, number of needle passes, volume of injectate, and accidental dural puncture were all significant risk factors for minor hemorrhagic complications.	Authors concluded that epidural steroid injection is safe in patients receiving aspirin-like antiplatelet medications. Minor worsening of neurologic function may occur after epidural steroid injection and must be differentiated from etiologies requiring interventions.	The results of this study strongly show that discontinuation of antiplatelet therapy and continued therapy with Aspirin was not essential and is not associated with major hemorrhagic complications.
Lagerkranser et al 2017 (52,53)	Authors in these 2 manuscripts studied neuraxial blocks and spinal hematoma with review of 166 case reports published from 1994 to 2015 with descriptions of demographics, risk factors, diagnosis, treatment, and outcomes. They utilized extensive search criteria in various languages across the globe. They also compared the previous reports published in 1992, 1994, and 1996 (175-177) 37% of patients with hematoma were not receiving antithrombotic therapy.	They identified 166 case reports on spinal hematoma after central neuraxial blockade during the years between 1994 and 2015. The annual number of case reports published during this period almost tripled compared with the 2 preceding decades. Authors identified 21 cases of hematoma from epidural injections with 17 after steroid injections, 5 in cervical, 4 in thoracic, and 8 in lumbar regions. They also identified 4 after percutaneous application of spinal cord stimulators. The authors identified spinal disease, which was identified as the most common of all spinal diseases in 14 cases with spinal disease reported in 37 cases of 166 cases.	Authors concluded that anti-hemostatic drugs, heparins in particular, are still major risk factors for developing spinal bleeding. Other risk factors were hemostatic and spinal disorders and complicated blocks, especially bloody taps, whereas multiple attempts did not seem to increase the risk of bleeding. They recommended that suspicion of spinal hematoma calls for the consultation of surgeon without delay. MRI was the recommended diagnostic tool. Surgical evacuation within 12 hours from the sign of motor dysfunction seems to lead to the best outcome, even though many patients operated as late as after more than 24 hours did regain full motor function.	This report is an extensive review of epidural hematoma of all origins, specifically of epidural injections for chronic pain with a prevalence of 21 cases of hematomas and 3 cases hematoma after spinal cord stimulation. Hematomas were identified in 37% of patients without antithrombotic therapy. Significant information is provided in this review indicating the risk of bloody taps, and prompt surgical intervention to improve outcomes. Limitations include lack of assessment after appropriate cessation of antithrombotic therapy.

NSAIDs = nonsteroidal anti-inflammatory drugs; SCS = spinal cord stimulator; MRI = magnetic resonance imaging

deaths 5 strokes, one pulmonary embolism, and one myocardial infarction in patients when anticoagulants were stopped; however, they have not reported any bleeding complications in patients where anticoagulants were continued, although, with inclusion of a small number of epidural injections with high risk. Kumar et al (178) reported a case of pulmonary embolism after discontinuation of warfarin during spinal cord stimulation trial. Linn et al (179) also reported right middle cerebral artery infarction with persistent left hemiparesis, neglect and dysarthria with L5-S1 epidural steroid injection after discontinuation of warfarin for 9 days preprocedure. Manchikanti et al (138) in providing 2 case reports and literature review described 2 cases of thromboembolic complications with cessation of antithrombotic therapy. Table 5 shows reported thromboembolic and cardiovascular complications related to discontinuation of antiplatelet or anticoagulation therapy.

3.4 Case Reports of Bleeding Complications

There were 35 case reports discovered citing epidural hematoma in patients with or without continued antithrombotic therapy during an interventional technique (130-138,148,180-204).

Table 6 shows reports of bleeding and epidural hematoma associated with interventional techniques in patients without antithrombotic therapy (136,138,140,144,180,183,185,190,192-195,197, 201,203,204,206,208,209), whereas Table 7 shows reports of bleeding in patients with discontinued antithrombotic therapy (130,133-135,138,139,148,181,187-189,191).

Table 8 shows reports of bleeding complications and epidural hematoma in patients with continuation of antiplatelet therapy with interventional techniques. Multiple reports of bleeding complications with continuation of anticoagulant therapy during interventional techniques were identified (131-133,136,137,182,199,200,202).

Table 4. Interlaminar epidural procedures with continuation or discontinuation of antithrombotic therapy.

	_	Aspirii	ı	Ţ	Varfari	n	Cl	opidog	rel	Aspi	rin + C	Others		lostazo ioxapa	
	D	C	T	D	C	T	D	С	T	D	С	T	D	C	T
Manchikanti et al (32)															
Cervical Epidural	161	249	410	12	1	13	67	10	77	34	18	52	NA	NA	NA
Thoracic Epidural	19	30	49	1	1	2	7	1	8	NA	5	5	NA	NA	NA
Lumbar Epidural	63	128	191	23	NA	23	16	14	30	28	9	37	NA	NA	NA
Percutaneous Adhesiolysis	68	148	216	24	NA	24	16	21	37	37	32	69	NA	NA	NA
Epidurals - Total	311	555	866	60	2	62	106	46	152	99	64	163	NA	NA	NA
Endres et al (42)															
Lumbar interlaminar	NA	NA	NA	169	25	194	69	15	84	NA	NA	NA	0	2	2
Epidurals - Total	311	555	866	229	27	256	175	61	236	99	64	163	0	2	2

None = Not on antithrombotic drugs; D = discontinued; C = continued; T = total

Others: Sacroiliac joint interventions, occipital nerve blocks, intercostal nerve blocks, stellate ganglion block, or lumbar sympathetic blocks

Table 5. Thromboembolic and cardiovascular complications related to discontinuation of antiplatelet or anticoagulation therapy.

Study	Type of study	Complications
Endres et al (42)	Observational report of interventional techniques	 2 patients died 5 suffered strokes 1 suffered pulmonary embolism 1 suffered myocardial infarction
Kumar et al (182)	Case report of dorsal column stimulator trial	Pulmonary embolism without lasting complications
Linn et al (183)	Case report of L5/S1 epidural steroid injection	Right middle cerebral artery infarction with persistent left hemiparesis, neglect, and dysarthria
Manchikanti et al (31)	Online survey	Reports of epidural hematoma: 55 Reports of thromboembolic complications: 162
Manchikanti et al (138)	Case report and literature review of interventional techniques	• 2 cases of thromboembolic complications with cessation of antithrombotic therapy.

Table 6. Reports of bleeding and epidural hematoma associated without antithrombotic therapy with interventional techniques.

Study/Year	Case Report	Conclusions
Swicegood et al, 2017 (136)	Authors reported a cervical epidural hematoma in a 41-year-old Caucasian female with controlled hypertension, but with no drug therapy contributing to bleeding. She received uneventful epidural injection and developed neurological dysfunction necessitating surgical decompression. She recovered without residual dysfunction.	Epidural hematoma developed in a patient without antithrombotic or anticoagulant therapy. Authors described this case of epidural hematoma in a patient with no risk factors or anticoagulant or antiplatelet therapy.
Manchikanti et al, 2017 (138)	Authors reported a case of 61-year-old female without risk factors or anticoagulant therapy. Epidural injection was performed in the cervical spine without difficulty. Patient developed symptomatology leading to the diagnosis of cervical epidural hematoma within 3 hours after the procedure undergoing surgical intervention due to progressive neurological dysfunction.	Epidural hematoma occurred after required period to discontinue aspirin. This patient had no other risk factors or other anticoagulant therapy. Patient was also successfully managed conservatively without surgical intervention. Patient developed epidural hematoma without any risk factors or anticoagulant therapy.
Manchikanti et al, 2018 (140)	Cervical epidural hematoma was reported in a healthy 43-year-old functioning female patient without overt risk factors for procedural hemorrhagic complications including baseline coagulopathy, baseline dysfunction, and intake of anticoagulants. There were no technical difficulties in performing the procedure. Patient developed symptomatology leading to the diagnosis of cervical and thoracic epidural hematoma with progressive neurological dysfunction necessitating surgical decompression.	Authors concluded that epidural hematoma may happen in cervical epidural injections considered as high risk procedure without antiplatelet or anticoagulant therapy and also without any risk factors.
Berrigan et al, 2018 (144)	Cervical epidural hematoma was reported in a 62-year-old healthy female patient after dry needling in cervical and thoracic musculature. Patient developed neurological dysfunction leading to the diagnosis of cervical and thoracic epidural hematoma confirmed by MRI. Due to stable neurological function, surgical intervention was deferred. She was managed with conservative management without residual dysfunction.	Authors report an epidural hematoma with dry needling, which is considered as extremely unusual in a patient without risk factors or antiplatelet or anticoagulant therapy.
Desai & Dua, 2014 (180)	This is a case report of a 72-year-old female undergoing right transforaminal epidural injection at L3 and L4. Four days after the injection patient reported progressive right lower extremity weakness, worsening sensory loss, and ambulatory dysfunction. An MRI with gadolinium enhancement showed focal abnormal signal with involvement of the right L4-5 neuroforamina that extended slightly far laterally, consistent with a small hematoma, affecting L4 nerve root. There was no history of coagulopathy.	This case report shows that though extremely rare, perineural or foraminal hematoma may occur as a serious complication of transforaminal epidural injection, even in the setting of a standardized procedure.
Ghaly, 2001 (183)	A 56-year old man who developed Brown-Séquard syndrome from a spinal epidural hematoma after fluoroscopically-guided cervical steroid injection. The patient reported immediate sharp shooting pain in the upper extremities on introduction of epidural Tuohy needle. Within half an hour, a neurological deficit occurred at C7/8 and right Brown-Séquard syndrome developed. MRI showed C6 to T2 spinal epidural hematoma with cord compression. Emergency spinal bilateral decompressive laminectomies and evacuation of spinal hematoma were performed within an expected delay of 10 hours from the onset of neurological deficit from C6 to T12.	Patient developed epidural hematoma without any risk factors or anticoagulant therapy.
Stoll & Sanchez, 2002 (185)	A healthy 34-year old man with no evidence of coagulopathy and not taking antiplatelet medication suddenly had onset of acute cervical myelopathy from a large cervical epidural hematoma 8 days after cervical epidural steroid injection. The patient developed a Brown-Séquard type of myelopathy manifesting by severe weakness of the left arm and leg and right-sided numbness and loss of temperature appreciation. The diagnosis was made by CT scan. Following prompt surgical evaluation of the clot, the patient made a near complete recovery.	The authors concluded that this case illustrated that epidural hematoma may occur in the absence of known risk factors. The delayed onset and the absence of risk factors have implications for the use of epidural steroid injection in chronic pain management.
Lee et al, 2011 (190)	A 58-year old woman presented with quadriparesis and neck pain after dry needling. MRI of the spine revealed a hyperintense mass in the T2 weighted at C2-T2 level, which proved to be an epidural hematoma. The diagnosis was made with MRI and decompression was carried out. The patient recovered completely. There were no risk factors and the patient was not on antiplatelet therapy.	The authors concluded the spinal hematoma caused by dry needling in this case was probably the result of unintentional needling of the spinal canal, similar to that caused by central neural blockade.
Ozdemir et al, 2007 (193)	Epidural steroid injection was performed with a #18 gauge Tuohy needle which revealed a dural puncture in a 40-year old male. The second attempt at L3/4 level was successful. MRI showed a right frontal subdural hematoma. His headache was relieved after strict bedrest, intravenous hydration, and analgesics. The patient was discharged with full recovery after one week without neurological dysfunction.	The authors concluded that intracranial subdural hematoma for accidental dural puncture during epidural steroid injection is a rare complication, even though there were no risk factors in this patient.

Table 6 (cont.). Reports of bleeding and epidural hematoma associated without antithrombotic therapy with interventional techniques.

Study/Year	Case Report	Conclusions
Lee et al, 2007 (195)	A healthy 38-year-old woman underwent a series of right transforaminal epidural injections at the C7-T1 level. Approximately 4 days after the final injection, she awakened with severe upper thoracic pain and progressive loss of sensation in the lower extremities. MRI of the spine showed a large hematoma extending from T1 to T5. The patient underwent surgical decompression. Within 6 months patient regained full strength and sensation in both lower extremities.	The authors cautioned that the physician should be aware that symptoms from a slowly developing epidural hematoma can present even days after an injection.
Domenicucci et al, 2017 (197)	A report of acute spinal epidural hematoma after acupuncture extending from C2 through T12. Due to rapid improvement with the patients' neurological symptoms, conservative treatment was adopted with excellent long-term results.	Acupuncture may produce epidural hematoma.
Keane et al, 1993 (201)	A case report of a 65-year-old female with development of epidural hematoma at T5-T6 several hours after acupuncture. Conservative management resulted in complete recovery.	Acupuncture may produce epidural hematoma.
Kim & Park, 2015 (203)	A 39-year-old female underwent cervical epidural injection at C6 under fluoroscopic guidance. Five days after epidural injection she complained of mild headache that was increasing in the upright position with a normal CT scan. At this time, she was treated with conservative management for post dural puncture headache. One month later, she presented with a severe headache that was not relieved by analgesic medication, which changed in character from being positional to non-positional during the preceding month. Brain magnetic resonance imaging revealed a chronic subdural hematoma along the left convexity. Emergency Burr Hole drainage was performed and the headache abated.	This report indicated that an intracranial chronic subdural hematoma presenting with intractable headache after cervical epidural steroid injection should be considered a possible complication. In addition, the event of an intractable and changing post dural puncture headache after epidural injection suggests further evaluation for diagnosis of an intracranial hematoma.
Takawira et al, 2012 (204)	A 52-year old male patient underwent spinal cord trial stimulation placement from T8 through T10 levels bilaterally with epidural entry at T12 and L1 interspace. 72 hours after the placement with good pain relief, he suddenly noticed relocation of stimulation to his right flank, with an abrupt onset of 10/10 burning lower back pain radiating down both lateral thighs and accompanied by inability to lift his knees. The patient also exhibited neurological deficits and loss of rectal tone. The trial leads were immediately removed and an MRI was performed. This showed an epidural or subdural fluid collection surrounding and compressing the thecal sac and spinal cord, extending from C7 to L3, which was interpreted as possibly a rapidly developing epidural hematoma. The epidural hematoma was a "thin spread" which was postulated as secondary to dislocation of the lead rather than trauma of the insertion. Epidural hematoma resolved rapidly and no surgical intervention was required with full recovery.	This is another case of spinal cord trial stimulation causing epidural hematoma with full recovery, but may cause severe problems. The authors noted that based on the retrospective analysis using a Manufacturer and User Facility Device Experience (MAUDE) database, the incidence of epidural hematoma was about 0.19%.
Smith et al, 2010 (206)	The authors reported 2 cases of epidural hematoma out of 4 complications related to spinal cord stimulations admitted to an acute spinal cord rehabilitation hospital over a 4-month period. The first patient with hematoma after placement of stimulator trial leads developed acute new mid back pain associated with rapid progressive weakness and motor loss to his bilateral lower extremities on day 5 associated with urinary retention. The pattern of weakness began in the right lower extremity and then progressed to the left lower extremity. A thoracic CT myelogram and CT of the thoracic spine with intrathecal contrast showed an epidural hematoma which extended 2 levels above the area of the spinal cord stimulator placement. The patient was not on any anticoagulants and all the studies were normal. On postoperative day 5, the patient underwent a thoracic T8/9 laminectomy with excision of epidural hematoma and removal of spinal cord stimulator electrodes. The patient developed permanent spinal cord injury	Spinal cord hematoma developing after a trial without any risk factors. Recovery was poor after acute rehabilitation.
Smith et al, 2010 (206)	The authors reported 2 cases of epidural hematoma out of 4 complications related to spinal cord stimulations admitted to an acute spinal cord rehabilitation hospital over a 4-month period. The authors reported a second patient, a 66-year old female with a history of chronic low back pain with stimulator leads placed in a patient with moderate to severe canal stenosis at L1/2 and L2/3 with epidural hematoma and air spanning T9-L2/3 and mass effect increasing the canal narrowing of the L1/2 and L2/3 levels. She underwent decompressive thoracic and lumbar laminectomies of T11-L2 levels. After acute inpatient rehabilitation stay, she recovered full motor strength in bilateral lower extremities. However, she was independent with limited community ambulation and had full bladder and bowel function recovery. She was diagnosed with spinal cord injury at T7.	Spinal cord hematoma developing after trial in a patient with severe spinal stenosis; however, without any antithrombotic therapy with moderate recovery after acute rehabilitation.

Table 6 (cont.). Reports of bleeding and epidural hematoma associated without antithrombotic therapy with interventional techniques.

Study/Year	Case Report	Conclusions
Chen et al, 1997 (208)	A case of a 48-year-old female developing epidural hematoma one week after acupuncture at L1-2. Patient was managed conservatively with complete recovery.	Acupuncture may produce epidural hematoma.
Eftekhar et al, 2005 (192)	A 74-year-old male patient developed epidural hematoma 2 days after acupuncture at L2-3. Patient underwent laminectomy with complete recovery.	Acupuncture may produce epidural hematoma.
Chen et al, 2006 (194)	A 30-year-old male patient with upper back pain developed epidural hematoma from C7-T3 one hour after acupuncture. Patient underwent laminectomy with complete recovery.	Acupuncture may produce epidural hematoma.
Nam et al, 2010 (209)	A 60-year-old male patient with back pain developed epidural hematoma 5 days after acupuncture from L4-S1. Patient underwent laminectomy with complete recovery.	Acupuncture may produce epidural hematoma.

 $\label{thm:continuation} \begin{tabular}{l} Table 7. \textit{Reports of bleeding and epidural hematoma in patients after discontinuation of antithrombotic therapy after interventional techniques. \end{tabular}$

Study	Antithrombotic/ Anticoagulant	Number of Days Stopped Prior to Procedure	Case Report	Conclusion
Benyamin et al (130)	Clopidogrel	12 days	Acute epidural hematoma formation was reported in cervical spine after interlaminar epidural steroid injection despite discontinuation of clopidogrel for 12 days. Procedure was performed atraumatically between C7 and T1 under fluoroscopic guidance and contrast injection. The patient started complaining of severe pain immediately after transfer to the recovery area. The patient underwent an expedited cervical spine MRI identifying a large epidural hematoma which was surgically decompressed with full recovery. Patient was given 30 mg of ketorolac intramuscularly for pain.	Epidural hematoma occurred in a patient after stopping of clopidogrel 12 days prior to the procedure. Ketorolac may contribute to exacerbation of hematoma. However, the patient was already significantly symptomatic prior to administration of ketorolac.
Giberson et al (133)	Low dose aspirin 81 mg and other NSAIDs	One week	This is a case report of a 70-year old patient undergoing spinal cord stimulator trial lead placement. He discontinued low dose aspirin 81 mg and other NSAIDs one week before the trial. After the removal of the trial after 4 days, the patient developed acute onset of burning lower thoracic pain and lower extremity weakness and spasms. An emergent MRI image revealed epidural hematoma extending from T8 to L3 with significant cord compression. The patient underwent an emergent T8 to T11 laminectomy and evacuation of the hematoma. He had complete resolution of his symptoms and was discharged home in good condition.	The authors reported, along with this case, another case where the patient took aspirin the day of the removal of the leads and developed hematoma. Consequently, they concluded aspirin must be discontinued. However, it is puzzling that this patient was on aspirin; however, he discontinued 7 days prior to the procedure.
Page et al (134)	Warfarin	Discontinued 7 days earlier with preoperative INR of 1.0	This case report involves a 67-year old woman undergoing interlaminar epidural injection with significant spinal stenosis at multiple levels undergoing interlaminar epidural injection with an #18 gauge Tuohy needle utilizing a paramedian approach with a single attempt. The patient developed weakness in legs the next day, 18 hours after the procedure. She had minimal relief with pain after the procedure. That evening around 11:30 pm, she was taken to the emergency room with weakness and inability to completely empty the bladder with an emergent MRI, which showed epidural hematoma from L3 to L5 with significant compression of the cauda equina. The patient underwent laminectomy within 4 hours of arrival to the emergency room. Total time from injection to the operating room was approximately 35 hours. She developed a permanent foot drop.	The authors described presence of spinal stenosis and the interlaminar approach seem to be significant risk factors in this patient. The epidural hematoma developed despite discontinuation of warfarin 7 days before the surgery.
Swicegood et al (135)	Aspirin, 81 mg	7 days	Authors reported a thoracic epidural hematoma after interlaminar epidural injection in a patient on 81 mg aspirin therapy which was discontinued 7 days prior. Epidural injection was performed at T10-11 leading to symptomatology developing of neurological dysfunction within 2 hours. Surgical decompression was carried out with rapid recovery.	Epidural hematoma developed despite stopping low dose aspirin 81 mg within the required time of 7 days.

 $\label{thm:cont.} \begin{tabular}{l} Table 7 (cont.). Reports of bleeding and epidural hematoma in patients after discontinuation of antithrombotic therapy after interventional techniques. \end{tabular}$

Study	Antithrombotic/ Anticoagulant	Number of Days Stopped Prior to Procedure	Case Report	Conclusion
Manchikanti et al (138)	Aspirin	9 days	A case of cervical epidural hematoma in a 68-year-old Caucasian female patient was presented. Patient was on 81 mg of aspirin which was discontinued 9 days prior to the procedure. Following an uneventful cervical interlaminar epidural injection she developed symptomology with the diagnosis leading to epidural hematoma confirmed by MRI. She was managed conservatively with improving symptomatology and recovered completely.	Epidural hematoma occurred after required period to discontinue aspirin. This patient had no other risk factors or other anticoagulant therapy. Patient was also successfully managed conservatively without surgical intervention.
Kim et al (139)	Clopidogrel, aspirin, and beraprost	7 days	Authors reported a lumbar epidural hematoma with late onset, 3 weeks after the performance of epidural steroid injection in a patient with spinal stenosis at L4-5 interspace. Patient was on clopidogrel, aspirin, and beraprost following infrarenal abdominal aortic aneurysm repair. Clopidogrel and aspirin were stopped 7 days prior to the epidural injection. Symptomatology developed 3 weeks after the epidural injection with formation of a large hematoma to cause severe thecal compression. Patient was managed conservatively without residual complications.	Epidural hematoma with late onset 3 weeks after the procedure in a patient after withholding appropriately clopidogrel and aspirin for 7 days. The authors also identified spinal stenosis as a contributing risk factor.
Caputo et al (148)	Dabigatran	7 days	A 70-year old patient underwent lumbar interlaminar epidural injection under CT guidance at L4/5. Dabigatran was started 24 hours after the injection. The patient had complete resolution of the symptoms. 48 hours after the injection, the patient started developing numbness in both lower extremities and had nearly complete paraplegia of the bilateral lower extremities. An emergent MRI showed an acute stenotic lesion, which was not present on the previous MRI. The patient was immediately decompressed. The patient was treated with multiple units of packed red blood cells prior to decompression. On surgical exposure, there was a large hematoma from L3-L5. Surgical evacuation of the hematoma was carried out with full neurological recovery 6 months after surgery.	The authors concluded that there is no reversal protocol for dabigatran. Dabigatran was stopped 7 days prior, which is longer than recommended duration. However, it was started one day after the procedure, which seems to have resulted in hematoma formation.
Shanthanna & Park et al (181)	Warfarin	Stopped for 4 days with an INR of 1.2 on the day of epidural injection	A 65-year-old male patient was treated with lumbar epidural injection at L3-4 in the sitting position with a second attempt which was atraumatic. Patient reported severe back pain and inability to stand 20 minutes after the procedure. An MRI scan performed within 3 hours of the patient's symptomatology revealed a localized dorsal/lateral thoracic epidural hematoma at T10 to T12. After neurosurgical referral, conservative management was provided with significant improvement within a day without having sustained any neurological deficit. An MRI done at the follow-up visit, a week later, showed a resolving hematoma.	Epidural hematoma developed after appropriate discontinuation of Warfarin and ideal INR of 1.2 within 20 minutes after the epidural injection. Epidural injection was at L3-4 level; however, hematoma was at T10 to T12. Authors concluded that the situation calls for optimization of all the relevant patient factors including anticoagulant therapy, use of appropriate technique, and the appropriate imaging modalities. However, spinal stenosis has been described as a significant risk factor to develop epidural hematoma with epidural injections.

 $Table\ 7\ (cont.).\ Reports\ of\ bleeding\ and\ epidural\ hematoma\ in\ patients\ after\ discontinuation\ of\ antithrombotic\ therapy\ after\ interventional\ techniques.$

Study	Antithrombotic/ Anticoagulant	Number of Days Stopped Prior to Procedure	Case Report	Conclusion
Chiravuri et al (187)	Clopidogrel	10 days	This is a case report in a 49-year-old male for placement of spinal cord stimulation trial leads for chest pain. Leads were attempted to be placed at L1-L2 with Tuohy needle under live fluoroscopic guidance contacting the L2 lamina with loss of resistance technique to the air. Subarachnoid puncture was noted with clear fluid flowing freely from the Tuohy needle. The needle was withdrawn into the epidural space and a spinal lead advanced to mid C7. Post operatively patient reported a positional headache accompanied by nausea and vomiting. This progressively worsened over the course of the day, changing in character with loss of positional component accompanied by emesis. His CT scan of the head showed a large subdural hematoma. Neurosurgical consultation was carried out and the patient underwent emergency craniotomy. The remaining post-surgical course was uneventful; however, on the postoperative day 3 patient recalled falling at home one day prior to the spinal cord stimulator implant, striking his head without loss of consciousness.	Even though this case presents acute intracranial subdural hematoma secondary to unintentional dural puncture during placement of permanent spinal cord stimulator lead, it also raises questions in reference to the causal relationship of the procedure and development as there was history of a fall. Even then, there is need for careful follow-up of patients with a known post dural tear. Failure to identify uncommon adverse events in patients with complicated spinal cord stimulator implantation may lead to permanent injury.
Xu et al (188)	Warfarin, Aspirin, Enoxaparin	6 days and 24 hours before procedure	Warfarin and aspirin stopped 6 days before the procedure with bridging over to enoxaparin twice daily, with the last enoxaparin dose at least 24 hours before the procedure. A 78-year old woman underwent lumbar interlaminar epidural injection under fluoroscopic guidance. 30 hours after the 3rd lumbar interlaminar epidural injection, the following morning after the 2nd shot of enoxaparin, the patient developed excruciating radicular leg pain. CT scan of the lumbar spine was inconclusive. Subsequently, MR imaging of the lumbar spine showed a lumbar epidural hematoma that spanned from L2 to L5 causing severe thecal sac compression. She was treated with bilateral lumbar laminectomies, foraminotomies and hematoma evacuation from L2-L5. The patient had unremarkable postoperative course. The epidural hematoma developed despite cessation of all drugs within the guidelines and restarting of enoxaparin within the guidelines.	Despite strict adherence to anticoagulation guidelines, epidural hematoma developed.
Ain et al (189)	Warfarin, Enoxaparin	Warfarin was stopped for 6 days and Enoxaparin was stopped for over 24 hours prior to the injection. INR was 1.2	A patient with renal insufficiency received an epidural injection at L3-L4 interspace. Enoxaparin regimen was reinstituted 24 hours after the procedure. 48 hours after the procedure she reported 100 times worse back pain in the same location as before the epidural injection. Magnetic resonance imaging study showed severe central canal stenosis centered at L3-4 from a posterior epidural hematoma. Epidural hematoma extended from L2 through L4. Her anticoagulation regimen was discontinued and managed conservatively. The second day the patient experienced increased numbness and weakness in the lower extremities. Subsequent MRI revealed increasing size of the epidural hematoma extending from L1 to L5. Decompression laminectomies were performed from L2 to L4 and dexamethasone was given for 3 days.	Patient developed epidural hematoma after epidural injection despite strict adherence to the guidelines for neuraxial anesthesia and anticoagulation regarding administration of low molecular weight heparin.
Reitman and Watters (191)	Aspirin (325 mg)	7 days	A 62-year-old woman underwent an uncomplicated cervical epidural steroid injection. She developed acute onset of axial pain followed by progressive quadriparesis within a matter of 8 hours. Emergency CT scan suggested posterior cord displacement consistent with an anterior spinal hematoma from C3 through C5. She was taken to the operating room and appropriate decompression was carried out showing anterior subdural hematoma. Patient continued to be quadriplegic, even though rapidly gained full function in the left upper and lower extremities. She developed acute meningitis and cardiac arrests and finally died.	Epidural hematoma can occur even after appropriate cessation of aspirin. Despite rapid diagnosis and surgical intervention, the outcome was bad in this case.

NSAIDs = non-steroidal anti-inflammatory drugs; MRI = magnetic resonance imaging; INR – international normalized ratio; CT – computed tomography

Table 8. Reports assessing bleeding complications and epidural hematoma in patients with continuation of antiplatelet therapy or drugs potentially or increase bleeding with interventional techniques.

	/otto-demonstration A		
Study	Anticoagulant	Case Report	Conclusions
Buvanendran et al (131)	Aspirin	71 year old woman with post-laminectomy syndrome presented for SCS trial. Patient had multiple medical problems and was taking Aspirin 81mg daily until the procedure day. Two 14-gauge introducer needles were placed with a single atraumatic pass via a paramedian approach toward the T12 to L1 interspace. 2 cylindrical octet electrodes were advanced into the dorsal epidural space with the aid of fluoroscopy. Next morning patient experienced severe bilateral lower extremity pain so the leads were removed and she subsequently developed lower extremity weakness. Emergent MRI showed T2-T10 epidural hematoma and she underwent surgery within 7 hours of symptoms onset. Patient did well post-operatively and regained her baseline functions.	ASRA guidelines may not be applied towards SCS leads placement and aspirin should be held 8-10 days prior implantation of SCS electrodes.
Chien et al (132)	Ketoralac, fluoxetine, fish oil, and vitamin E	A 66 year-old woman with chronic renal insufficiency and neck pain due to multi-level neuroforaminal stenosis and degenerative intervertebral discs and on Ketoralac, fluoxetine, fish oil, and vitamin E was treated with epidural injection. Utilizing a loss of resistance to saline technique, an 18-gauge Tuohy-type needle entered the epidural space at C6-7. After negative aspiration, 4 mL of saline with 80 mg of methyl-prednisolone was injected. Immediately thereafter, the patient reported significant spasmodic-type localized neck pain without any neurologic status changes and it was treated with 30mg ketorolac injection. Later on, she developed a sudden onset of acute tetraplegia. An emergent MRI in ED demonstrated an epidural hematoma extending from C5 to 17 and coagulation studies were normal. She underwent a bilateral C5-T6 laminectomy with epidural hematoma evacuation and was discharged to an acute inpatient rehabilitation hospital. She had permanent lower extremity paralysis with bladder dysfunction.	Combined antiplatelet effects of ketorolac, fluoxetine, fish oil, and vitamin E may have played a role in compromised hemostasis in this this patient. Using a smaller blunt needle may also decrease the incidence of epidural hematoma.
Giberson et al (133)	Excedrin (acetaminophen, aspirin, and caffeine) taken on the morning of removal of the trial	This is a case report of a 53-year old man with trial leads placed with good pain relief. The patient had a successful trial and removal of trial leads without any problems except for there was some bleeding at the exit site, which stopped with local pressure. At this time, the patient reported that he took Excedrin the morning before the removal of the trial leads. The patient reported immediate difficulty, which was diagnosed as epidural hematoma from T5-L2 with spinal cord compression. The patient was treated with surgical intervention 2 days after admission with development of permanent weakness in his left leg. It appears the patient has taken aspirin that morning. The patient took aspirin on the same day when the leads were removed. Time to peak effect of aspirin is less than 1 hour with plasma half-life of 30 minutes (36).	The authors concluded that aspirin taken on the day of insertion of the leads or removal of the leads can be dangerous.
Swicegood et al (136)	Aspirin	Author reported a case of cervical epidural hematoma in a 76-year-old Caucasian male patient receiving low dose aspirin, 81 mg, and Vitamin D3. Aspirin was continued. Patient also had multiple arthritic changes in the cervical spine with moderate canal stenosis and bilateral foraminal narrowing. Patient developed symptomatology leading to the diagnosis of epidural hematoma which required surgical decompression. Patient recovered without residual dysfunction; however, patient was also administered with 30 mg of intravenous ketorolac which may have contributed to the epidural hematoma.	Authors describe that epidural hematoma may occur with low dose aspirin therapy; however, authors also emphasized on the importance of risk factors related to the anatomical factors.
Jenkie et al (137)	Fish oil	A 49-year-old woman with a history of moderate spinal stenosis in the cervical area underwent cervical epidural injection. Two years prior, she had an anterior cervical disc fusion at C5-C6 for significant C6 radiculopathy and had subsequently developed disc herniations above and below the level of fusion. After the procedure and the development of symptoms, a cervical MRI showed a 7 mm epidural hemorrhage extending superiorly to the C2-C3 level and inferiorly to the T4 level. The majority of the hemorrhage was at the C6-C7 level associated with moderate cervical stenosis due to a cervical disc herniation at that same level. She underwent emergent surgical decompression and evacuation of the hematoma through multiple laminectomies at C6, C7, and T1. The patient had immediate relief of her symptoms and regained her wrist strength.	The authors concluded the need to evaluate fish oil as a predisposition to patients with bleeding complications when used in higher doses.
Williams et al (182)	Indomethacin	A diagnosis of cervical epidural hematoma was made and was confirmed by CT scan after a seventh epidural at <i>C7/</i> T1 in several years in a patient who was taking indomethacin. Immediate surgery revealed an epidural hematoma. During the operation blood results became available showing that INR was 1.0, the clotting time was 41 seconds with a control of 40 seconds, and the platelet count was within normal limits. Bleeding time was not measured. His recovery was complicated with wound breakdown requiring a skin graft and urinary retention which required prostatectomy. Subsequently he made a full recovery.	Authors concluded that repeated epidural steroid injections should preferably be performed in centers equipped to accurately to diagnose and promptly treat this rare but serious complication.

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Table 8. (cont) Reports assessing bleeding complications and epidural hematoma in patients with continuation of antiplatelet therapy or drugs potentially or increase bleeding with interventional techniques.

Study	Antithrombotic/ Anticoagulant	Case Report	Conclusions
Maier et al (199)	Ticlopidine and clopidogrel	Two cases were presented with lumbar sympathetic blockade in 2 patients under medication with irreversible platelet aggregation inhibitors. The first patient was a 71-year-old male patient with intermittent daudication and a trophic leg ulcer due to progressive peripheral arterial disease. He was on ticlopidine 500 mg/day for stroke prevention in the presence of carotid artery stenosis. Patient underwent left-sided lumbar sympathetic block under fluoroscopy. There was no known vascular puncture. Two days later, a widespread skin hematoma was recognized. Four days later (6 days after the first block), a second block was performed under fluoroscopic visualization which involved intravascular needle penetration. That is. Following this, the patient developed a large retroperitional hematoma. He was managed conservatively. The second case involved a 79-year-old female patient with generalized peripheral arterial disease, polyneuropathy, coronary artery sclerosis, and a history of stroke with treatment with 75 mg per day of clopidogrel. Clopidogrel was discontinued 3 days prior to lumbar sympathetic blocks were performed at 1.3 level with attempts with no intravascular injection. She complained of burning groin and medial thigh pain. Her pain was improved with opioid. She was walking on the ward without complaints. An hour later she was found pulseless and resuscitation attempts were unsuccessful. The autopsy revealed a massive coagulated hematoma beneath the left psoas muscle with enlarged retroperitoneal hemorrhage. There were no visible puncture lesions in larger vessels.	The authors concluded that irreversible platelet inhibitors should be discontinued for at least 7 days prior to any invasive technique, and any hemorrhage is not controllable by external compression. They also concluded that in case of high thrombotic risk, patients should be switched to a different therapeutic regimen with heparin up to 6 hours prior to blockade. They believe that groin pain and, particularly, pain at the medial side of the thigh, seem to be early signs of bleeding complications of lumbar sympathetic block.
(200) (200)	Clopidogrel, possibly Aspirin	This case report describes a patient with acute onset of quadriparesis admitted to a teaching hospital with a large cervical epidural hematoma, developing numbness and weakness of his arms and legs within 30 minutes after the performance of the cervical epidural steroid injection. At the time of injection, he was taking clopidogrel, possibly aspirin, and delofenear that had been started after a prior coronary anjoplasty. He also had a prior uneventful cervical epidural steroid injection 11 days before the incident in question, but it is not clear if he was taking these antiplatelet drugs then. His Pf, INR, and PPT were within normal limits. His platelet count was appropriate. The last doses of antiplatelet drugs were unclear. He underwent an emergency C3-T3 laminectomy and evacuation of cervical epidural hematoma. Approximately 14 hours after the epidural injection postoperatively upper extremity strength in his upper extremities, but the paralysis of his lower extremities remained.	The authors of this case report cautioned that these drugs inhibit platelet adhesion to the vascular endothelium and shear stress-induced platelet aggregation and section. After discontinuation of clopidogrel therapy, platelet aggregation and bleeding time returned to baseline within 5 days. Authors stated that there has been no case report of spinal hematoma after neuraxial block in patients receiving clopidogrel; however, there was a case report with ticlopidine. They recommended that neuraxial blocks be postponed for 5 - 7 days in patients who are receiving several antiplatelet drugs.
Choi et al (202)	Cilostazol	This is a case report of a patient undergoing caudal epidural injection for spinal stenosis on cilostazol. Three days after the caudal injection, the patient developed severe burning pain radiating into both hips and difficulty standing. The patient's symptoms started 6 hours after the caudal injection and worsened with time. An MRI showed epidural hematoma with acute cord compression at L2 through S1 with concomitant central canal compromise, severe at L2/3 and L3/4 levels. Emergency decompressive laminectomy and evacuation of the hematoma was performed uneventfully over 4 hours. Patient made full recovery.	This is probably the first case report of epidural hematoma following a caudal epidural injection. Multiple guidelines have recommended that Cilostazol need not be stopped prior to the spinal procedures. Spinal stenosis has been reported as a major risk factor; however, this procedure was performed caudally avoiding such risk. Cilostazol, combined with aspirin, did not prolong bleeding time above significant increase of bleeding time caused by aspirin alone or dopidogrel.

ASRA = American Society of Regional Anesthesia; MRI = magnetic resonance imaging; INR = international normalized ratio; PT = prothrombin time; PPT = partial thromboplastin time

Table 9 shows summary reports of a number of epidural hematomas with epidural injections, acupuncture and dry needling, and spinal cord stimulation lead placement in patients without antithrombotic therapy, with antithrombotic therapy withheld for appropriate duration, and with antithrombotic therapy continued (Figs. 2-4). A total of 38 epidural hematomas were described in 35 case reports. There were 20 in cervical spine, 8 in thoracic spine, and 10 in lumbar spine. Of these, 19 patients were on no antiplatelet therapy, 11

patients had their antithrombotic therapy discontinued, and 8 continued antithrombotic therapy. Further, as shown in Fig. 4, epidural injections were responsible for 25 cases of hematomas, 7 cases were secondary to acupuncture or dry needling and 6 were related to spinal cord stimulation. There was one case report of caudal epidural injection with cilostazol (202), and in one case report we were unable to obtain full manuscript (207). Cases of chronic subdural hematoma and cases of abdominal hematomas were not included.

Table 9. Reports of epidural hematomas.

	No Anticoagulant Therapy	Anticoagulant Discontinued	Anticoagulant Continued	Total
Cervical Epidural	8	Aspirin – 2	Aspirin – 1	
Stimulator trial	0	Clopidogrel – 1	Fish oil – 2	
Acupuncture/Dry Needling	3	Warfarin – 1	Indomethacin – 1	
			Clopidogrel – 1	
TOTAL	11	4	5	20
Thoracic Epidural	0	Aspirin 1		
Stimulator trial	3	Aspirin 1		
Acupuncture/Dry Needling	1	_	Aspirin 2	
TOTAL	4	2	2	8
Lumbar Epidural	1	Aspirin + Clopidogrel - 1,	Cilostazol (caudal) 1	
Stimulator trial	0	Warfarin – 3		
Acupuncture/Dry Needling	3	Dabigatran – 1		
TOTAL	4	5	1	10
COMBINED TOTALS	19	11	8	38

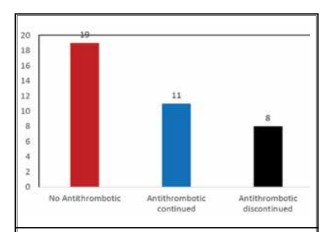


Fig. 2. Epidural hematoma incidence based presence or absence of anticoagulant/antiplatelet therapy.

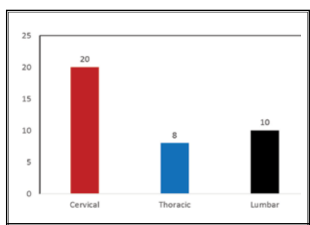


Fig. 3. Epidural hematoma incidence based on spinal regions.

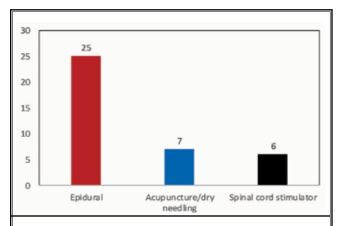


Fig. 4. Epidural hematoma cases reported based on type of procedure.

4.0 GUIDELINES AND RECOMMENDATIONS

Guidelines and recommendations were based on risk stratification, pharmacology of antithrombotics and anticoagulants, and application of available evidence.

4.1 Risk Stratification

Interventional techniques performed in the spine and other regions for chronic cancer and noncancer pain patients face variable risks depending on anticoagulant or antithrombotic therapy, age, anatomy, the specific region of interest, and obesity. Consequently, various authors have described procedural classification according to the potential risk for serious bleeding.

Raj et al (212) stratified risk score based on technique related bleeding risk and patient-related bleeding risk factors. This risk classification took various factors into consideration including a sharp or blunt needle, use of fluoroscopy and lack of fluoroscopy.

Breivik et al (36) in a comprehensive topical review of reducing risk of spinal hematoma from spinal epidural and pain procedures based their recommendations on extensive review of 166 case reports published since 1994 through 2015 (52,53), pharmacology of drugs, and available clinical evidence relating to complications whether or not the antithrombotics were continued or discontinued.

Narouze et al (40) provided guidance for interventional spine and pain procedures in patients on antiplatelet and anticoagulant medications. They provided extensive review of the literature, considered pharmacology, and current literature with development of risk stratification.

Deer et al (39) provided recommendations on bleeding and coagulation management in neurostimulation devices. They also provided bleeding risk stratifications

for neuromodulation procedures. Their classification showed spinal cord stimulation trial and implant, dorsal root ganglion stimulation, and intrathecal catheter and pump implant as high to intermediate risk neuromodulation procedures. Further, they classified deep brain stimulation and motor cord stimulation as high-risk neuromodulation procedures. This is in contrast to other guidance where spinal cord stimulation trial and implant, dorsal root ganglion stimulation, and intrathecal catheter and pump implant are considered as high risk.

Oprea et al (20) published risk stratification, perioperative, and periprocedural management of patients receiving anticoagulant therapy based on bleeding risk for each procedure, pharmacology, and evidence of risk of development of bleeding complications as well as thromboembolic risks.

Lagerkranser (52) and Lagerkranser and Lindquist (53) published extensive review of neuraxial blocks and spinal hematoma with review of 100 case reports published from 1994 to 2015 with demographics and risk factors, as well as diagnosis, treatment, and outcome. They concluded that the annual number of published cases of spinal hematoma after central neuraxial blocks increased during 1994 to 2015 compared to previous years. Case reports on elderly women accounted for this increase; however, they also emphasized that anti-hemostatic drugs, heparins in particular, still continue to be major risk factors for developing post central neuraxial blockade spinal bleedings. They also described other risk factors related to hemostasis and spinal disorders and complicated blocks, especially spinal stenosis and "bloody taps," whereas multiple attempts do not seem to increase the risk of bleeding (Table 10). Further, they conceded that in a large number of cases, no risk factor was reported.

Lagerkranser (52) also identified patient-related risk factors as shown in Table 10. Topping the list of the risk factors is spinal disease in general and spinal stenosis in particular. Surprisingly, INR of \geq 1.5 was the least related risk factor second only to liver disease.

Among the drug-related risk factors, these were the most common risk factors with the intake of antihemostatic drugs in 105 of 166 cases (63%), and with no drug therapy in 37% of cases. Further, in 47 of the cases, more than one anti-hemostatic drug was implicated in 12 cases in 3 such drugs. Table 11 shows drug related risk factors with low molecular heparin topping with 31% of the cases, aspirin at third level

with 16% of cases, vitamin K antagonist with 11% of the cases, NSAIDs also at 11%, and adenosine diphosphate (ADP)-receptor inhibitors and phosphodiesterase inhibitors at 2.5% and 1.3% respectively.

They also concluded that guidelines issued around the turn of the century do not seem to have affected the number of published reports. They also concluded that the complications continue to occur despite following the appropriate guidelines and strict adherence to the recommendations. Consequently, they concluded that there are no guarantees against such bleedings to occur, even if guidelines are strictly followed.

Lagerkranser (52) and Lagerkranser and Lindquist (53) also concluded that suspicion of a spinal hematoma calls for the consultation of an orthopedic or neurological surgeon without delay. Surgical evacuation within 12 hours from the first sign of motor dysfunction seems to lead to the best outcome, even though many patients operated on later than 24 hours did regain full motor function. They also concluded that despite the poor prognosis after surgical evacuation of spinal hematoma, the outcomes after central neural blockade in general have improved significantly over time.

Our data analysis in this report showed interlaminar epidural injections, spinal cord trial lead placement through interlaminar approach, and spinal stenosis as risk factors. Overall, as shown in Fig. 2, when antithrombotics were continued, epidural hematomas occurred in 8 patients compared to 19 with no antithrombotic therapy and 11 with antithrombotic therapy appropriately discontinued. This provides a ratio of 27% incidence of hematomas when antithrombotics were continued compared to 73% with either no antithrombotics, or with appropriate discontinuation of antithrombotics. There were larger number of patients in the group where antithrombotics were discontinued with 11 compared to 8 in continued group. The majority of the hematomas were reported in the cervical region with 20, followed by 10 in the lumbar region and 8 in thoracic region (Fig. 3). Further analysis also showed that 25 of 38 cases were related to epidural injections, 7 were related to either acupuncture or dry needling, and 6 were related to spinal cord stimulation (Fig. 4). Thus, epidural injection, specifically in the cervical spine, is considered as high risk, along with spinal cord lead placement. This is illustrated in Figs. 5 to 7. As shown in Fig. 5, cervical epidural hematoma incidence was highest in patients without any antithrombotic therapy with 11 patients, followed by continuation of antithrombotic therapy in 5 patients, followed by in 4 patients where antithrom-

Table 10. Patient related risk factors as described in review of 166 cases of spinal hematoma.

Disorder	No. of cases
Spinal stenosis	14
Spinal tumor	6
Ankylosing spondylitis	4
Osteoporosis	4
Herniated disc	2
Scoliosis	2
Spondylosis	2
Spina bifida occulta	1
Dural AV-fistula	1
Spinal AVM	1
Spinal disease (total)	37
Thrombocytopeniaa	13 ^b
Renal insufficiency	10
INR ≥ 1.5	6°
Liver disease	4

^a Platelet count $< 150 \times 10$ 9 /L.

Source: Lagerkranser M. Neuraxial blocks and spinal haematoma: Review of 166 case reports published 1994-2015. Part 1: Demographics and risk-factors. *Scand J Pain* 2017; 15:118-129 (52).

Table 11. Drug classes reported, in order of frequency in 160 cases of spinal hematoma.

	No. of cases (percentage)
Low molecular weight heparin	50 (31)
Unfractionated heparin	39 (24)
Acetylsalicylic acid	25 (16)
Vitamin-K antagonists	17 (11)
Non-steroidal anti-inflammatory drugs	17 (11)
Dextran	6 (3.8)
Fibrinolytics	4 (2.5)
ADP receptor inhibitors	4 (2.5)
Phosphodiesterase inhibitor (dipyridamole)	2 (1.3)
Direct fXa inhibitor (rivaroxaban)	2 (1.3)
Direct thrombin inhibitor (dabigatran)	1 (0.6)
Fondaparinux	1 (0.6)
Selective serotonin receptor inhibitor	1 (0.6)

Source: Lagerkranser M. Neuraxial blocks and spinal haematoma: Review of 166 case reports published 1994-2015. Part 1: Demographics and risk-factors. *Scand J Pain* 2017; 15:118-129 (52).

^b Including 2 obstetric cases, one with HELLP (haemolysis, elevated liver enzymes, and low platelets), and one with eclampsia, and one patient with idiopathic thrombocytopenic purpura, discovered postlaminectomy.

^c In one case without anti-vitamin K treatment.

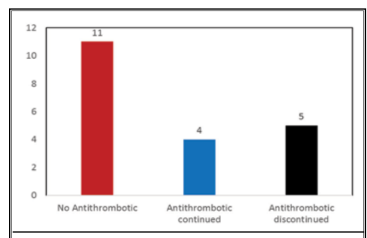


Fig. 5. Cervical epidural hematoma incidence bbased presence or absence of anticoagulant/antiplatelet therapy.

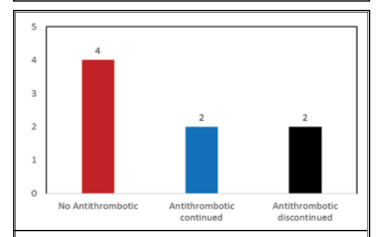


Fig. 6. Thoracic epidural hematoma incidence based presence or absence of anticoagulant/antiplatelet therapy.

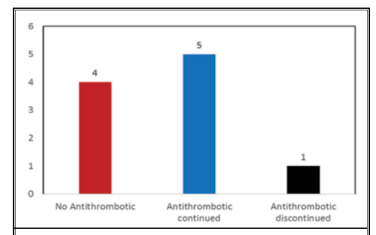


Fig. 7. Lumbar epidural hematoma incidence based presence or absence of anticoagulant/antiplatelet therapy.

botic therapy was discontinued. Overall, in the cervical spine, in 67% of the patients developing hematoma formation, there were either no antithrombotics or antithrombotics were discontinued appropriately. Results were even more striking in lumbar spine as shown in Fig. 7. Overall, there were 10 instances of lumbar epidural hematoma, 4 in patients without antithrombotic therapy, 5 with appropriate discontinuation and only one patient where it was continued with cilostazol which is rarely utilized and none of the guidelines recommend it to be discontinued. Thus, in the lumbar spine, only one patient or 10% of epidural hematomas were related to continuation of antithrombotic therapy. In contrast, in the thoracic spine the results were similar to the cervical spine with 6 patients developing hematoma, either without anticoagulant therapy or after appropriate discontinuation (4+2) and in 2 patients with continuation of antithrombotic therapy with a 25% incidence of epidural hematoma in the thoracic spine in patients continuing antithrombotic therapy similar to cervical spine and higher than lumbar spine.

The previous studies have shown low molecular weight heparin as the most frequently used drug followed by unfractionated heparin, followed by aspirin (Table 12). The present assessment showed aspirin as the most common drug with a total of 7 of 38 patients; however, it was continued only in 3 patients; whereas it was discontinued in 4 patients. Warfarin was also common with or without enoxaparin; however, it was discontinued according to the guidelines. Clopidogrel was reported in 3 cases with discontinuation in 2 patients and continuation in one patient. Many of the commonly utilized antithrombotics have not been identified in the present analysis.

Multiple factors and the available literature in reference to the adverse consequences of anticoagulant and antithrombotic therapy with interventional techniques were utilized in developing risk stratification. The following classification for interventional techniques has been developed (Table 13) (1,20,31-40,52,53,198-200,212). The classification describes low risk, intermediate risk, and

high-risk procedures. However, based on comorbid medical conditions and other risk factors of coagulopathies and concurrent use of other anticoagulants and antiplatelets, the procedural risk classification may be changed from low risk to intermediate, intermediate to high risk.

Of all the low risk procedures as shown in Table 13, caudal epidural injection was the only one which has reported an epidural hematoma with continuation of cilostazol (202). This is extremely unusual and probably coincidental. Consequently, with caudal epidural injections and other low risk procedures, antithrombotic and anticoagulant therapy may be continued with the appropriate guidelines.

Intermediate risk procedures include multiple procedures performed, constituting a great proportion of procedural prevalence. This risk stratification is somewhat different from the one proposed by ASRA (40). Justifiably we have included caudal percutaneous adhesiolysis into intermediate risk procedures as there have not been any case reports. Further, lumbar transforaminal epidural injections at L4, L5, S1, and sympathetic blocks have been included in these categories due to one or 2 case reports. Lumbar interlaminar epidural injections performed between L5 and S1 are included in intermediate risk procedures; whereas, procedures performed at L4-5 and at higher levels are included in high risk procedures. A majority

of the lumbar epidural hematomas developed despite discontinuation of antithrombotic therapy and these were performed above L5-S1 interspace. If epidural hematoma develops at L5-S1 it will have significantly higher space availability to be asymptomatic and to be managed conservatively.

The high risk procedures include the majority of the procedures performed in the cervical and thoracic spine. These guidelines are in contrast to ASRA guidelines with the inclusion of cervical, thoracic, and high lumbar above L5-S1 due to the available literature and potential issues related to epidural hematoma requiring surgical exploration and spinal cord damage.

Table 12. Drug classes reported in order of frequency.

	Continued	Discontinued	Total
None			19
Aspirin discontinued	4	3	7
Warfarin discontinued	0	4	4
Clopidogrel	1	2	3
Fish oil	2	0	2
Indomethacin	1	0	1
Cilostazol	1	0	1
Dabigatran	0	1	1

Table 13. Classification of interventional techniques based on the potential risk for bleeding.

Low-Risk Procedures	Intermediate-Risk Procedures*	High-Risk Procedures*
1. Trigger point and muscular injections (including piriformis injection) 2. Peripheral joints 3. Peripheral nerve blocks 4. Sacroiliac joint and ligament injections and nerve blocks 5. Caudal epidural injections 6. Ganglion impar blocks	1. Facet joint interventions (intraarticular injections, nerve blocks and radiofrequency neurotomy) 2. Lumbar transforaminal epidural injections at L4, L5, S1 3. Lumbar intradiscal procedures 4. Hypogastric plexus blocks 5. Lumbar sympathetic blocks 6. Peripheral nerve stimulation trial and implant 7. Pocket revision and implantable pulse regenerator/intrathecal pump replacement 8. Caudal percutaneous adhesiolysis 9. Lumbar percutaneous disc decompression (L4/5 or below) 10. Lumbar vertebral augmentation (below L4) 11. Intervertebral spinous prosthesis 12. Lumbar discography	1. Cervical, thoracic, and high lumbar (above L4-L5) interlaminar epidurals 2. Cervical, thoracic and lumbar above L3 transforaminal epidural injections 3. Spinal cord stimulator trial and implant 4. Percutaneous adhesiolysis with interlaminar or transforaminal approach 5. Percutaneous disc decompression (above L4/5) 6. Sympathetic blocks (stellate ganglion; thoracic splanchnic, celiac plexus) 7. Thoracic and cervical intradiscal procedures 8. Vertebral augmentation, lumbar (above L4), thoracic and cervical 9. Intrathecal catheter and pump implant 10. Interspinous prosthesis and MILD*
	13. Lumbar interlaminar epidural injections at L5-S1	

^{*}Patients with high risk of bleeding (e.g., old age, history of bleeding tendency, concurrent uses of other anticoagulants/antiplatelets, liver cirrhosis or advanced liver disease, and advanced renal disease) undergoing low or intermediate-risk procedures should be treated as intermediate or high risk, respectively.

4.2 Pharmacologic Aspects and Hemostasis Monitoring

The main categories of antithrombotics and anticoagulants are described as; platelet inhibitors, interfering with platelet aggregation (clumping) and thrombus formation; anticoagulants interfering with formation of the clotting, thereby reducing fibrin formation and preventing clots from forming and expanding; and fibrinolytics interfering with the final clot.

Monitoring of hemostasis is performed with multiple standard tests including platelet count, activated partial thromboplastin time (APTTa), and INR. The normal platelet count is x 109 L-1 from 150 to 350 with recommended counts of greater than 50 for vital lifethreatening indication, greater than 80 for strong indication, and greater than 100 for weak indication (36). A normal INR is considered as 0.9 to 1.2. Consequently, for epidural injections, INR of less than 1.5 is ideal for high risk and moderate risk procedures, and less than 1.8-2.00 is ideal for low risk procedures.

Additional advanced hemostatic tests for monitoring of hemostasis include multiple viscoelastic tests like thromboelastography or thromboelastrometry extensively used to evaluate liver disease (213). However, this parameter has been studied for safe epidural catheter removal with the conclusion that the tests were not well validated in this context and there were frequent false negative test results (214,215). Even then, a clearly abnormal curve indicates deranged hemostasis and must be taken seriously (215,216).

4.2.1 NSAIDS and Aspirin

NSAIDs inhibit cyclooxygenase enzymes COX1 and COX2, which inhibit prostaglandin production to decrease the inflammatory response. Thus, NSAIDs have analgesic effects and are used for minimizing pain. Thromboxane A2 is produced via COX1 enzyme activity, which is a potent thrombus activator. Aspirin is an irreversible inhibitor of COX1 and has significant clinical benefits for preventing thrombus formation. In response to aspirin, more prostacyclin is produced by endothelial cells, but there is no additional thromboxane made as there are no nuclei in platelets, thus, there is a greater percent of prostacyclin to thromboxane, thinning the blood. Elevated bleeding risk is a concern for a small portion of patients, but adverse effects are rare. Prostacyclin (PGI2) synthesis from vascular endothelial cells is dependent on COX2 and has anti-platelet effects. High doses of aspirin reduce PGI2 production which can abolish the anti-platelet effect of low dose aspirin.

Low-dose aspirin anti-platelet effects last for 7-10 days, as bone marrow directed platelet renewal is required for clotting to resume. Low-dose aspirin therapy is well established to reduce the risk of cardiovascular events in patients with acute coronary syndromes, cerebral infarct, or occlusive vascular disease (217). However, recently published large scale evidence (15-19) shows lack of benefit with aspirin for primary prevention of cardiovascular events, while it does increasethe risk of bleeding.

The data reported by Lagerkranser et al (52) and the data derived from the present analysis are shown in Tables 11 and 12. Aspirin was the most common drug of the 19 cases of epidural hematoma reported in this series with 7 with use of aspirin. Four patients discontinued aspirin appropriately and 3 patients continued. In the data by Lagerkranser et al (52), aspirin was utilized in 16% of patients reporting spinal hematoma of 160 cases constituting a total of 18% in the present series. Eight percent continued aspirin and 11% discontinued aspirin. The most common drug reported in Lagerkranser et al (52) series was low molecular weight heparin in 31% of the cases and Warfarin in 11% of the cases. In the present series, Warfarin with or without low molecular weight heparin was utilized in 4 patients with 11% utilization.

Table 14 shows pharmacokinetic and pharmacodynamic characteristics of aspirin and NSAIDs. Time to 50% recovery of platelet function with aspirin is shown as 3 days (36). Antiplatelet function of irreversible inhibitors is dependent mainly on platelet regeneration than drug half-life. While bleeding complications have been reported without any drug therapy, majority of the bleeding complications were related to aspirin in 18% while 11% discontinued and 7% continued.

For other NSAIDs, unlike aspirin, the platelet effects of these drugs are directly related to systemic plasma drug concentrations and influenced by the pharmacokinetic clearance of these medications. It takes approximately 5 half-lives for systematic elimination. Recent recommendations (40) have revised recommendations for aspirin, whether to continue or discontinue in the perioperative period for interventional pain procedures. Decision-making is based on the reason for aspirin utilization, a multitude of risk factors including vascular anatomy surrounding the target area, degree of the invasiveness of the procedure, and potential sequelae associated with perioperative bleeding. Thus, aspirin for primary prophylaxis can be stopped without any hesitation. The major consideration in withholding

Table 14. Characteristics of aspirin and NSAIDs.

	Aspirin or acetylsalicylic acid (Oral Low Dose)	NSAIDs	
Target	COX -1 irreversible	COX-1 reversible, COX-2	
Time to peak effect	0.5 hours	Varies	
Plasma Half Life	0.5 hours	Variable from 1 to 72 hours	
Renal elimination	+	+	
Time to 50% recovery of platelet function	3 days	1 day	
Hours to C-Max	0.5 hours	~0.5 hours	
Metabolism	Hepatic	Hepatic	
Bioavailability	60%	50-95%	
Antihemostatic effect	++	+	

 $NSAID = non-steroidal \ anti-inflammatory \ drug; \ COX = cyclo-oxygenase$ Rating of antihemostatic effect and renal elimination: (+) = insignificant; + = low; ++ = moderate; +++ = pronounced; ++++ = high.

aspirin is the thromboembolic risk. Based on the available evidence, it appears that aspirin discontinuation for 4 days may be sufficient. In contrast, for non-steroid anti-inflammatory agents, recommended discontinuation by ASRA is one day for diclofenac, ibuprofen, and Ketoralac. Recommended discontinuation time is 2 days for etodolac and indomethacin. Discontinuation is about 4 days for meloxicam and Naprosyn, 6 days for nabumetone, and 10 days for piroxicam and oxaprozin. However, the evidence for stopping non-steroidal anti-inflammatory agents other than aspirin seems to be very limited. Stopping these drugs may become a practical issue and patients may not like it to stop all the drugs and complain of significantly more pain. Consequently, based on available, very limited, evidence, the clinician may continue or discontinue.

4.2.2 ADP Receptor Inhibitors

ADP-receptor inhibitors inhibit platelet aggregation. The drugs in this category utilized for clinical applications include Clopidogrel (Plavix®), Prasugrel (Effient®), Ticlopidine (Ticlid®), and Ticagrelor (Brilinta®). Table 15 shows comparative pharmacokinetics and pharmacodynamics of ADP-receptor inhibitors.

Clopidogrel is the prototypical thienopyridine drug that inhibits the P2Y12 receptor. The P2Y12 receptor is activated by ADP binding and promotes platelet aggregation. Depending on the dosage schedule, the maximal platelet aggregation inhibitory effects of clopidogrel are reached within 3 to 7 days. After discontinuation, recovery of platelet inhibition occurs 50% after 3 days and full recovery after one week (218) Eighty percent of subjects demonstrated normal platelet aggregation by the 4th day (218). Other studies have demonstrated the

recovery of platelet function after cessation of aspirin in volunteers in surgical patients after 3 days in volunteers and within 4-6 days in surgical patients (211). In this study, by day 6, all of the subjects had restored platelet aggregation to at least 85% of the baseline level.

Ticlopidine also belongs to the thienopyridine group and is maximally aggregated after 8 to 11 days of a 500 mg per day dosage schedule. After withdrawal of 72 hours, there is still a lingering effect as there is an irreversible inhibition of platelet function (219). Prasugrel acts by antagonizing ADP at the platelet's purine receptors, and aggregation is thus noncompetitively and irreversibly inhibited.

Prasugrel, or Effient, has significantly higher irreversible antiplatelet activity compared to clopidogrel with time to peak effect of one hour. Thus, administration of first dose results in around half of the platelets being inhibited within the first hour of taking this medication. Following three to five days of therapy, the steady-state inhibition of platelet aggregation reaches around 70% (220). As a prodrug, prasugrel is rapidly metabolized to active and inactive metabolites. These metabolites have varying elimination rates, although the active metabolites have an elimination half-life of 7 hours, with a wide range of 2-15 hours (221).

Lastly, a distinct ADP-receptor inhibitor is Ticagrelor, which directly inhibits P2Y12 receptors (222-224). While Ticagrelor is metabolized to active metabolites, the original compound is responsible for the majority of the inhibitory effects (223,224). A notable advantage of Ticagrelor is rapid effect, with peak platelet inhibition after 2 to 4 hours of intake (225). These medications undergo hepatic conversion to active metabolites, which are then eliminated by the kidneys (226). In addition, glycoprotein

Table 15. Comparative pharmacokinetics/pharmacodynamics of ADP-receptor inhibitors.

	Clopidogrel (Plavix®)	Prasugrel (Effient®)	Ticlopidine (Ticlid®)	Ticagrelor (Brilinta®)
Target	P2Y12ADP	P2Y12ADP	P2Y12ADP, also inhibits liver CYP2C19 and CYP2B6	P2Y12ADP
Antithrombotic activity	++	+++	++	++++
Time to Cmax	3-7 days	3-5 days	8-11 days	2-4 hours
Time to peak effect	4 hours to 4 days	1 hour	3-5 days	2.5 hours
CYP metabolism	CYP1A2, CYP2B6, CYP2C9, CYP2C19 and CYP3A4/5	CYP450-mediated (primarily CYP3A4 and CYP2B6)	Cytochromes P450	CYP3A4
Bioavailability	> 50%	≥ 79%	> 80%	36%
Protein binding	94–98%	Active metabolite: ~98%	98%	>99.7%
Plasma half-life	7-8 hours (inactive metabolite)	~7 hours (range 2 hours to 15 hours)	12 hours (single dose) 4-5 days (repeated dose)	7 hours (ticagrelor), 8.5 hours (active metabolite AR-C124910XX)
Renal elimination	50% kidney, 46% biliary	Urine (~68% inactive metabolites); feces (27% inactive metabolites)	Renal and fecal	Biliary
Time to 50% recovery of platelet function	3 days	3 days	6 days	1.5 days

CYP = cytochrome P450

IIB/IIIA receptors are less activated, causing a reduction in fibrinogen fixation and platelet crosslinking.

ADP inhibitors were described as responsible for the formation of epidural hematoma after central neuraxial blockade in 2.5% of 160 cases or 4 cases. In the present assessment, clopidogrel either in combination with aspirin or ticlopidine was utilized in a total of 3 cases, with 2 cases being discontinued appropriately and one case it was continued. Thus, ADP inhibitors with reports available for only one or 2 drugs showed any relevance in 8% of the cases with only one case or 2.6% of cases where it was continued and in 5% of the cases it was discontinued. This is similar to the reports from Lagerkranser et al (52).

4.2.3 Phosphodiesterase (PDE) Inhibitors

Phosphodiesterase inhibitors include Cilostazol (Pletal®) and Dipyridamole (Persantine®). These medications selectively inhibit phosphodiesterase, which leads to an increase in intracellular cyclic adenosine monophosphate (cAMP) and subsequent reversible inhibition of platelet aggregation (227). Additionally, Dipyridamole blocks thromboxane synthase, the thromboxane receptor, and the cellular reuptake of adenosine into platelets, red blood cells, and endothelial cells. This results in increased adenosine in the extracellular space and inhibition of formation of cytokines and

proliferation of smooth muscle cells. Absorption of Dipyridamole occurs in the gastrointestinal tract and is pH dependent. Gastric acid suppressors and proton pump inhibitors inhibit absorption, which can be prevented via buffered additives added to the medication (228). An additional advantage of Cilostazol is inhibition of PDE3A, which is selective to vascular smooth muscle cells and results in vasodilatation. Cilostazol is administered at 100 mg twice daily and reaches maximum plasma levels after three hours. It is eliminated via hepatic metabolism and is excreted in the urine (229). Thus, cilostazol is contraindicated in those with severe renal insufficiency. For interventional procedures, phosphodiesterase inhibitors have been considered as safe to continue. However, risk may increase with addition of aspirin. Limited data exists evaluating the risk of perioperative surgical bleeding with cilostazol (40) and no standard perioperative guidelines are available (230). Further, if the medication is discontinued, at 50 hours (approximately 5 half-lives), less than 5% of the drug remains in plasma and improvements in platelet aggregation have been demonstrated, despite continuous prior dosing (210,231).

There is a single case report of bleeding complications associated with interventional technique (202). In this case report, the patient underwent a caudal epidural injection developing epidural hematoma extending from L2 through S1 with concomitant central canal compromise, severe at L2-3 and L3-4 levels. This patient received, in the past, lumbar interlaminar epidural injection without any side effects. Emergency decompression laminectomy was carried out with the patient making full recovery. This is the only case report in interventional techniques. Cilostazol alone without aspirin is recommended to be continued during interventional techniques. Lagerkranser et al (52) also reported 2 cases of phosphatase inhibitors with dipyridamole with an incidence of 1.3% in 160 cases. Overall cilostazol is considered as a low risk drug and its continuation is appropriate.

4.2.4 Glycoprotein GPIIb/IIIa Inhibitors

A final common component of platelet aggregation is the glycoprotein IIb/IIIa receptor. Specialized medications inhibit this receptor, potently inhibiting platelet aggregation while being reversible (172).

Abciximab (ReoPro®) is a Fab fragment of a humanized monoclonal antibody directed against the GFPIIb receptor. Abciximab inhibits over 80% of ADP-induced platelet aggregation and is given via IV administration. Additionally, thrombin generation is inhibited by Abciximab, which quickly binds to platelets with high affinity.

Eptifibatide (Integrilin®) is a cyclic peptide inhibitor of the fibrinogen binding site on the GPIIb receptor. Tirofiban (Aggrastat®) is an additional glycoprotein IIb/ IIIa receptor inhibitor, reaching maximum efficacy after 4 hours of administration, with 50-80% inhibition of platelet aggregation (232). Platelet function normalizes 8 to 24 h after stopping the IV infusion.

There were no case reports secondary to development of epidural hematoma in patients receiving glycoprotein IIB/IIIA. In addition, it appears that these drugs are not commonly used for prevention of thromboembolic activity.

4.2.5 Low-Molecular Weight Heparin

Low molecular weight heparins (LMWHs) inhibit the coagulation cascade via binding to antithrombin, which leads to a conformational change of antithrombin, which accelerates inhibition of factor Xa. LMWH has advantages: relatively high bioavailability, longer half-life, and ability for use once per day. Maximum efficacy levels are observed after 3-4 hours post subcutaneous administration, and elimination occurs after 4-6 hours in those with normal renal function (233). High molecular weight heparins (HMWH) catalyze the

inhibition of clotting factors IXa, Xa and thrombin by greatly enhancing antithrombin III activity, by causing a conformational change in ATIII exposing its reactive site. Testing is required to determine the dose effect on coagulation via partial thromboplastin time (PTT). HMWH is not absorbed by GI tract due to its large molecular weight, therefore IV or SC injection must be used. The short half-life of HMWH approximately 1h) means frequent injections or continuous infusion, and it is thus not considered suitable in an outpatient setting.

Low molecular weight heparin is one of the commonly used drugs and has been implicated in multiple cases of epidural hematoma. Lagerkranser et al (52) showed low molecular weight heparin being responsible for the highest number of cases in 31% of the reports. Our reports also show 2 cases of warfarin with bridging with LMWH being responsible for epidural hematoma (188,189), both in the lumbar spine.

4.2.6 Warfarin

Oral anticoagulants inhibit the synthesis of vitamin K-dependent clotting factors, which are factor II, VII, IX, and X. Warfarin blocks the gamma-carboxylation of glutamate residues in prothrombin and factors VII, IX, and X. This results in biologically inactive coagulation factor molecules. Vitamin K epoxide reductase is the enzyme that catalyzes the carboxylation reaction. Therapeutic doses of warfarin inhibit vitamin K epoxide reductase, which prevents the reductive metabolism of the inactive vitamin K epoxide to its active hydroquinone form. Synthesis is the primary target of oral anticoagulants (warfarin), therefore the effects of these medications are not apparent until previously-existing clotting factor turnover has occurred. Factor half-lives vary, from factor VII at 6-8 hours to factor II at 50-80 hours (234). Thus, it has a slow onset of action (8-12 hours) as existing clotting factors must be depleted, and the maximal effect occurs 3-5 days after administration. Warfarin is monitored by PT and INR, which is a normalized ratio of the patient's PT to that of a control sample (234). Age, female gender, and preexisting medical conditions such as hepatic, cardiac, and renal disease modify the patient's response to warfarin. Asian patients, for example, have higher sensitivity to warfarin and require lower doses than those patients of European descent (234). Dietary changes may alter the patient's clotting ability, and those on Warfarin are advised to avoid grapefruit and cranberry products, eat a consistent amount of leafy greens and other high vitamin K containing foods and are advised to limit herbal supplement intake of garlic, ginger, gingko biloba, ginseng, and fish oil. Warfarin may be reversed with administration of vitamin K, which is associated with multiple side effects.

Warfarin is one of the most common drugs utilized in patients undergoing interventional techniques. Multiple complications have been reported with case reports of epidural hematoma in patients with warfarin, despite if it was stopped as per the guidelines.

Lagerkranser et al (52) reported warfarin contributing to spinal hematoma in 11% of the cases. Our report showed a total of cases of warfarin which were stopped appropriately with 2 of them also receiving enoxaparin with a similar incidence of around 10%.

4.2.7 Direct Thrombin Inhibitors

Direct thrombin inhibitors include Dabigatran

(Pradaxa®), Argatroban (Acova™), Bivalirudin (Angiomax®), Lepirudin (Refludan®), Desirudin (IPRIVASK®), and Hirudin as shown in Table 16. Of all the direct thrombin inhibitors, Dabigatran may be reversed by Idarucizumab (Praxbind®), which was approved in 2015.

Dabigatran etexilate is an oral anticoagulant and is a prodrug that is converted to dabigatran in the plasma. After an oral dose, the peak effect is reached within 2 to four hours, and plasma half-life is 13 hours on average (235). Dabigatran dose recommendations depend on renal efficacy in the patient receiving the medication. In those with a creatinine clearance of greater than 30mL/minute, 150 mg is taken orally twice daily. For patients with lower creatinine clearance, 75mg twice daily is recommended. Dabigatran's function is via factor inhibition and not clotting factor depletion, thus, the administration of clotting factors is anticipated to be less effective in reversing the effects of dabigatran. Dabigatran

Table 16. Comparative pharmacokinetics/pharmacodynamics of direct thrombin inhibitor.

	Dabigatran (Pradaxa)	Argatroban (Acova)	Bivalirudin (Angiomax)	Lepirudin (Refludan)	Desirudin (IPRIVASK)	Hirudin
Target	Direct thrombin inhibitor	Direct thrombin inhibitor	Reversible direct thrombin inhibitor	Direct thrombin inhibitor	Direct thrombin inhibitor	Naturally occurring peptide anticoagulant
Time to Cmax	2-4 hours	1-3 hours	2 minutes	4 hours	1-3 hours	3 hours
Time to peak effect	0.5 – 2 hours	2 hours	15 min	0.5-2 hours	2 hours	3-4 hours
Metabolism	Metabolized via conjugation into 4 acyl glucuronides, not mediated by CYP450	СҮРЗА4	Proteolytic cleavage	Lepirudin is thought to be metabolized by release of amino acids via catabolic hydrolysis of the parent drug	Metabolized by stepwise degradation from the C-terminus possibly catalyzed by carboxypeptidase(s) such as carboxypeptidase A	Proteolytic cleavage
Bioavailability	3-7%(Oral)	100% IV	100% IV application only	100% (injection or infusion)	100%	100% IV
Protein binding	35%	54%	no	n/a	n/a	n/a
Plasma half-life	13 hours	50 minutes	~25 minutes in patients with normal renal function	1.3 hours	2-3 hours	80 minutes
Renal elimination	80% urine	Liver	Yes	Yes	Yes	Renal, about 48% (35% unchanged)
Linear PK	Yes	Yes	Yes	Yes	Yes	Yes
Time to 50% recovery of thrombin function	12 hours	2 hours	0.5 hours	1.5 hours	2 hours	2 hours
Reversal agents	Praxbind	NA	NA	NA	NA	NA

CYP = cytochrome P450; IV = intravenous

is mostly cleared by the kidneys. In those with normal kidney function, Dabigatran is excreted in 1-2 days post-discontinuation. This also depends on renal sufficiency of the patient taking the medication. There is one case report with epidural hematoma despite its discontinuation for 7 days prior to interventional techniques (152). Lagerkranser et al (52) also reported on a case of spinal hematoma out of 160 cases.

Argatroban is a small molecule direct thrombin inhibitor that is administered intravenously. It reaches steady-state plasma concentrations in 1-3 hours and is metabolized via the liver. It has a half-life of 50 minutes and is monitored by PTT. As it is metabolized hepatically, it is a viable alternative for Dabigatran, which is metabolized renally (236).

Bivalirudin works by binding specifically to the catalytic site, in addition to the anion-binding exosite of circulating and clot-bound thrombin. Bivalirudin is cleared by the kidney and thus is dose-dependent on overall renal function. It has as a half-life of 25 minutes in those with normal renal function, but this may be doubled in those with severe renal insufficiency (237).

Desirudin is a subcutaneously administered direct thrombin inhibitor and is indicated for the prevention of venous thromboembolism after total joint replacement. It is recommended that dosage adjustment and aPTT be monitored in patients with moderate-to-severe renal impairment. After intravenous administration, desirudin is removed rapidly via the renal system, with 90% of the dose removed from the plasma within two

hours. Plasma concentrations decline with a mean half-life of 2-3 hours. Subcutaneous administration demonstrates a half-life of 2 hours (238).

Hirudin has specific activity on fibrinogen and binds to and inhibits only activated thrombin, making it an extremely potent direct thrombin inhibitor. Thus, hirudin dissolves the formation of clots and thrombi and has therapeutic value in coagulation disorders. It is also able to act on complexed thrombin and does not alter other serum protein function or activity (239). Hirudin has a half-life of 2-3 hours and is monitored by aPTT, allowing close titration over a wide range of anticoagulative clinical desires. Activated clotting time (ACT) and prothrombin time (PT) are insensitive for monitoring hirudin .

4.2.8 Direct Factor Xa Inhibitors

Direct factor Xa inhibitors such as Rivaroxaban (Xarelto®) have been commonly used in the United States (Table 17). Of multiple Xa inhibitors available, apixaban (Eliquis) and rivaroxaban (Xarelto) can be reversed by Andexanet alfa (Andexxa®), a coagulation factor Xa (recombinant), which has been approved by the FDA for the urgent reversal of the anticoagulant effect in 2018 (240).

Rivaroxaban (Xarelto) has dual renal and hepatic clearance, with around one-third of the drug being active with each route of metabolism. This dual route of clearance makes accumulation less likely than other medications that are solely hepatically or renally

Table 17. Comp	aratıve pnarmacoкıı	1et1cs/pnarmacoayn	amics of airect jac	tor Aa innibitor.

	Apixaban (Eliquis)	Rivaroxaban (Xarelto)	Edoxaban (Savaysa, Lixiana)	Betrixaban (Bevyxxa)	Fondaparinux (Arixtra)
Target	Xa	Xa	Xa	Xa	Xa
Time to Cmax	1-3 hours	2-4 hours	1-2 hours	3-4 hours	2 hours
Time to peak effect	3-4 hours	2-4 hours	1-2 hours	3-4 hours	2-3 hours
CYP Metabolism	15%	32%	NR	NR	n/a
Bioavailability	66%	80%	>45%	34%	100%
Transporter	P-gp	P-gp/BCRP	P-gp	P-gp	P-gp
Protein binding	87%	>90%	55%	60%	94%
Plasma Half-life	8-15 hours	9-13 hours	8-10 hours	37 hours	17-21 hours
Renal elimination	25%	33%	35%	< 1%	100%
Linear PK	Yes	No	Yes	n/a	Yes
Time to 50% recovery of Xa	12 hours	12 hours	12 hours	19-27 hours	12 hours
Reversal AGENTS	Andexxa	Andexxa	NA	NA	NA

BCRP = breast cancer resistance protein; CYP = cytochrome P450; NR - not reported; P-gp = P-glycoprotein

cleared. Rivaroxaban is orally administered and has a half-life of 5.7 to 9.2 hours. Plasma protein binding of rivaroxaban is 92-95%. One third of the absorbed dose is excreted in the urine, and two-thirds of the dose is excreted as an inactive metabolite in the feces and urine. Rivaroxaban has the potential for drug interactions with medications that are P-glycoprotein inhibitors and those metabolized by CYP3A4 (241).

Apixaban (Eliquis®) is a specific factor Xa inhibitor like its counterpart, rivaroxaban. It is rapidly absorbed and reaches peak concentrations in 1-2 hours (242,243). Apixaban has an oral availability of 45% and has a relatively complex elimination pathway with both direct renal and intestinal excretion, with the latter being the majority (243,244). Edoxaban (Savaysa® or Lixiana®) was approved for prevention of venous thromboembolisms following lower limb orthopedic surgery in 2011 and is an oral direct factor Xa inhibitor that inhibits free factor A and prothrombinase activity. It has also been approved for the prevention of stroke and systemic embolism. Peak plasma concentrations are reached 1.5 hours after oral administration, and it has an elimination half-life of 10-14 hours when taken at 60mg once daily. It is excreted via both hepatic and renal systems (245). It is orally available, and not removed by dialysis.

Betrixaban (Bevyxxa®) is a potent oral factor Xa inhibitor that recently received FDA approval. It has exemplified promising results, as it has low hERG affinity and has reduced bleeding risk and prevented thromboembolism in clinical trials for orthopedic knee surgery (246-248). Betrixaban has the smallest percent of renal clearance, is INR/PTT insensitive, and has minimal liver metabolism. Another selective factor Xa inhibitor, Fondaparinux (Arixtra®) is 100% bioavailable and achieves maximum concentration in 1.7 hours of administration (249). Its extended half-life of 17 to 21 hours allows once-daily dosing (250).

There were no case reports in the present assessment; however, Lagerkranser et al (52) showed 2 cases of spinal hematoma with rivaroxaban yielding 1.3% prevalence among 160 cases developing spinal hematoma.

4.2.9 Thrombolytic Agents

Fibrinolysis is caused by thrombolytic agents via conversion of plasminogen and thrombi to plasmin in order to destroy clots. These "clot busters" such as recombinant tissue-type plasminogen activator (tPA), streptokinase, urokinase, tenecteplase, and reteplase are enzymes that have effects on both circulating and

tissue type plasminogen. The half-life of these thrombolytic drugs is generally a few hours, but the inhibition of plasminogen and fibrinogen may last for up to 27 hours after administration (251).

There were no reports of epidural hematoma development in patients receiving thrombolytic agents in our analysis; however, Lagerkranser et al (52) reported 4 cases or 2.5% in the review of 160 cases of spinal hematoma. Among the miscellaneous agents, Chien et al (132) reported a case of epidural hematoma in a patient receiving fluoxetine, fish oil, and vitamin E. There was also another case report by Jenkie et al (137) with fish oil and the development of cervical epidural hematoma leading to surgical intervention for decompression. Lagerkranser et al (52) also reported one patient on selective serotonin receptor inhibitor of 160 cases.

4.2.10 Herbal/Alternative Therapies

Garlic has a dose-dependent effect on bleeding, as it contains a compound called ajoene. Derived from allicin, the compound that provides garlic's flavor, ajoene inhibits granule release and fibrinogen binding and additionally inhibits aggregation of platelets via a variety of mechanisms. Prostacyclin, forskolin, indomethacin, and dipyridamole are all altered via ajoene's inhibition of granule release (252,253). Ginkgo Biloba has been used for thousands of years, and its mechanism is not entirely understood. Ginkgo is thought to antagonize platelet activating factor (PAF) and collagen leading to inhibition of platelet aggregation, resulting in several reports of spontaneous bleeding. Flavonol glycosides and terpene glycosides have been suggested to be the chemical compounds responsible for the increased bleeding events after intake of this medication (254). Ginseng is commonly used and reduces the effect of warfarin, declining peak INR levels. Ginsenosides are the major active ingredient of ginseng, and possibly induce cytochrome P450 enzymes to increase the metabolism of Warfarin and thus reduce its effect.

5.0 Recommendations

Based on the evidence from review of the current literature with inclusion of available studies (1), case reports (38), multiple guidelines and reviews (1,20,31-42,122-129) and consensus, the evidence appears as follows:

 There is good evidence for risk stratification by categorizing multiple interventional techniques into low-risk, moderate-risk, and high-risk. Also,

- their risk should be upgraded based on other risk factors.
- There is good evidence for the risk of thromboembolic events in patients who interrupt antithrombotic therapy.
- There is good evidence supporting discontinuation of low dose aspirin for high risk and moderate risk procedures for at least 3 days, and there is moderate evidence that these may be continued for low risk or some intermediate risk procedures.
- 4. There is good evidence that discontinuation of anticoagulant therapy with warfarin, heparin, dabigatran (Pradaxa®), argatroban (Acova®), bivalirudin (Angiomax®), lepirudin (Refludan®), desirudin (Iprivask®), hirudin, apixaban (Eliquis®), rivaroxaban (Xarelto®), edoxaban (Savaysa®, Lixiana®), Betrixaban(Bevyxxa®), fondaparinux (Arixtra®) prior to interventional techniques with individual consideration of pharmacokinetics and pharmacodynamics of the drugs and individual risk factors increases safety.
- There is good evidence that diagnosis of epidural hematoma is based on severe pain at the site of the injection, rapid neurological deterioration, and MRI with surgical decompression with progressive neurological dysfunction to avoid neurological sequelae.
- 6. There is good evidence that if thromboembolic risk is high, low molecular weight heparin bridge therapy can be instituted during cessation of the anticoagulant, and the low molecular weight heparin can be discontinued 24 hours before the pain procedure.
- 7. There is fair evidence that the risk of thromboembolic events is higher than that of epidural hematoma formation with the interruption of antiplatelet therapy preceding interventional techniques, though both risks are significant.
- 8. There is fair evidence that multiple variables including anatomic pathology with spinal stenosis and ankylosing spondylitis; high risk procedures and moderate risk procedures combined with anatomic risk factors; bleeding observed during the procedure, and multiple attempts during the procedures increase the risk for bleeding complications and epidural hematoma.
- There is fair evidence that discontinuation of phosphodiesterase inhibitors is optional (dipyridamole [Persantine], cilostazol [Pletal]. However, there is also fair evidence to discontinue Aggrenox [dipyri-

- damole plus aspirin]) 3 days prior to undergoing interventional techniques of moderate and high risk.
- 10. There is fair evidence to make shared decision making between the patient and the treating physicians with the treating physician and to consider all the appropriate risks associated with continuation or discontinuation of antithrombotic or anticoagulant therapy.
- There is fair evidence that if thromboembolic risk is high antithrombotic therapy may be resumed
 hours after the interventional procedure is performed.
- 12. There is limited evidence that discontinuation of antiplatelet therapy (clopidogrel [Plavix®], ticlopidine [Ticlid®], Ticagrelor [Brilinta®] and prasugrel [Effient®]) avoids complications of significant bleeding and epidural hematomas.
- 13. There is very limited evidence supporting the continuation or discontinuation of most NSAIDs, excluding aspirin, for 1 to 2 days and some 4 to 10 days, since these are utilized for pain management without cardiac or cerebral protective effect.

Table 18 shows guidelines for antithrombotic medication management during interventional spine procedures. This table also shows comparisons of ASIPP proposed guidelines and ASRA published guidelines. These recommendations differ from our previously published guidelines, as well as from ASRA guidelines based on the present analysis of the evidence.

General recommendations based on ASRA guidelines (40) are to discontinue the drugs for 5 half-lives; however, the exception to the 5 half-lives recommendation should occur in individuals with hepatic dysfunction, and renal dysfunction including nephrotic syndrome.

6.0 DISCUSSION AND LITERATURE REVIEW

6.1 Risk/Benefits of Antithrombotic Therapy

Cardiovascular disease in the United States contributes to the most mortality with 610,000 deaths per year contributing to one in every 4 deaths (58). Thrombotic complications contribute a significant proportion of cardiovascular related deaths. Various disorders contributing to thrombotic complications include atherosclerotic cardiovascular disease, defined as acute coronary syndrome, ischemic heart disease, peripheral arterial disease, atrial fibrillation, and multiple other

Table 18. Guidelines for antithrombotic medication management and spinal procedures (risk stratification described in Table 13).

Medication	Time to Wait After Last Dose of Medication Before Low Risk Interventional Techniques Are	r Last Dose fore Low Risk chniques Are	Time to Wait After Last Dose of Medication Before Moderate Risk Interventional Techniques Are	er Last Dose of e Moderate Risk chniques Are	Time to Wait After Last Dose of Medication Before High Risk Interventional Techniques Are	ast Dose e High Risk iiques Are	Timing of Therapy restoration or Restarting	apy restoration
	Pertormed ASTDD	ASD A (40)	Performed	4SD 4 (40)	Fertormed ACTDD	4SD 4 (40)	ASTED	ACD A (40)
	ASIFF	ASLA (40)	ASILI	(40) WSUA	ASIFF	ASRA (40)	ASILL	ASRA (40)
NSAIDS	May continue or	Stop 1-10 days	May continue or	Stop 1-10 days due	May continue or stop	Stop 1-10 days due	24 hours	24 hours
(COX 2)	stop 1-10 days due to lack of	due to lack of protective effect	stop 1-10 days due to lack of	to lack of protective effect	1-10 days due to lack of protective effect	to lack of protective effect		
	protective effect	4	protective effect		4			
Aspirin								
Low-Dose Aspirin	Continue or may stop for 3 days	Stop for 4 days	Continue or may stop for 3 days	Stop for 4 days	Stop for 5 days	Stop for 6 days	24 hours	24 hours
High Dose Aspirin	Continue or may stop for 3 days	Stop for 4 days	Continue or may stop for 3 days	Stop for 4 days	Stop for 5 days	Stop for 6 days	24 hours	24 hours
Antiplatelet Agents (Phosphodiesterase Inhibitors)	osphodiesterase Inhibi	tors)						
Dipyridamole (Persantine)	May continue	May continue	May continue	May continue	May continue or stop for 2 days	Stop for 2 days	12 hours	12 hours
Cilostazol (Pletal)	May continue	May continue	May continue	May continue	May continue or stop for 2 days	Stop for 2 days	12 hours	12 hours
Aggrenox (dipyridamole plus aspirin)	May continue	Stop for 4 days	May continue	Stop for 4 days	Stop for 5 days	Stop for 6 days	24 hours	24 hours
Platelet Aggregation Inhibitors	hibitors							
Clopidogrel (Plavix)	May continue	May continue	May continue or stop for 3 days	Stop for 7 days	Stop for 5 days	Stop for 7 days	12 hours	12 hours
Prasugrel (Effient)	May continue	May continue	May continue or stop for 6 days	Stop for 7-10 days	Stop for 6 days	Stop for 7-10 days	24 hours	24 hours
Ticlopidine (Ticlid)	May continue	NA	May continue or stop for 7 days	NA	Stop for 7-10 days	NA	24 hours	24 hours
Ticagrelor (Brilinta)	May continue	Continue	May continue or stop for 3 days	NA	Stop for 3-5 days	Stop for 5-10 days	24 hours	24 hours
Vitamin K Antagonists								
Warfarin	May stop for 2 days INR ≤ 3.0	INR < 3.0	Stop for 2-5 days INR ≤ 1.5	Stop for 5 days INR normalize	Stop for 2-5 days INR ≤ 1.5	Stop for 5 days INR normalize	24 hours	24 hours
Thrombin Inhibitors								
Dabigatran (Pradaxa)	May continue or stop for 2 days	May continue or stop for 2 days	Stop for 4-5 days 6 days - renal	Stop for 4-5 days 6 days - renal	Stop for 4-5 days 6 days - renal	Stop for 4-5 days 6 days - renal	24 hours	24 hours

Table 18 (cont.). Guidelines for antithrombotic medication management and spinal procedures (risk stratification described in Table 13).

Medication	Time to Wait After Last Dose of Medication Before Low Ris	r Last Dose ore Low Risk	Time to Wait After Last Dose of Medication Before Moderate Risk	er Last Dose of e Moderate Risk	Time to Wait After Last Dose of Medication Before High Risk	Last Dose e High Risk	Timing of Ther or Restarting	Timing of Therapy restoration or Restarting
	Interventional Techniques Are Performed	thniques Are	Interventional Techniques Are Performed	chniques Are	Interventional Techniques Are Performed	niques Are)	
	ASIPP	ASRA (40)	ASIPP	ASRA (40)	ASIPP	ASRA (40)	ASIPP	ASRA (40)
Anti-Xa Agents								
Apixaban (Eliquis)	May continue or stop for 2 days	May continue or stop for 2 days	Stop for 3-5 days	Stop for 3-5 days	Stop for 3-5 days	Stop for 3-5 days	24 hours	24 hours
Rivaroxaban (Xarelto)	May continue or stop for 1 day	May continue or stop for 1 day	Stop for 2 days	Stop for 3 days	Stop for 2 days	Stop for 3 days	24 hours	24 hours
Edoxaban (Savaysa, Lixiana)	May continue or stop for 1 day	NA	Stop for 3 days	NA	Stop for 3 days	NA	24 hours	24 hours
Heparins								
Heparin (treatment) - IV	Discontinue for 4 hours	Discontinue for 4 hours	Discontinue for 4 hours	Discontinue for 4 hours	Discontinue for 4 hours	Discontinue for 4 hours	24 hours	24 hours
Heparin (treatment) - SC	Discontinue for 8-10 hours	Discontinue for 8-10 hours	Discontinue for 8-10 hours	Discontinue for 8-10 hours	Discontinue for 8-10 hours	Discontinue for 8-10 hours	24 hours	24 hours
Low Molecular Weight Heparin	Discontinue for 24 hours	Discontinue for 24 hours	Discontinue for 24 hours	Discontinue for 24 hours	Discontinue for 24 hours	Discontinue for 24 hours	24 hours	24 hours
Thrombolytic Agents								
TPA, Streptokinase, Alteplase, Reteplase	May continue	May continue	Stop for 2 days	Stop for 2 days	Stop for 2 days	Stop for 2 days	24 hours	24 hours
GPIIb/IIIa Inhibitors								
Abciximab (ReoPro)	May continue	May continue	Stop for 1-2 days	Stop for 2-5 days	Stop for 1-2 days	Stop for 2-5 days	8-12 hours	8-12 hours
Eptifibatide (Integrilin)	May continue	May continue	Stop for 8 hours	Stop for 8-24 hours	Stop for 8 hours	Stop for 8-24 hours	8-12 hours	8-12 hours
Tirofiban (Aggrastat)	May continue	May continue	Stop for 8 hours	Stop for 8-24 hours	Stop for 8 hours	Stop for 8-24 hours	8-12 hours	8-12 hours
Miscellaneous								
Fondaparinux (Arixtra)	May continue	May continue	Stop for 4 days	Stop for 4 days	Stop for 4 days	Stop for 4 days	8-12 hours	8-12 hours

ASIPP = American Society of Interventional Pain Physicians; ASRA = American Society of Regional Anesthesia and Pain Medicine

ailments. These patients are widely prescribed with anticoagulants and antithrombotics in over 10% of the patients undergoing surgery or invasive procedures (255,256). However, a significant proportion of these patients may also undergo invasive procedures resulting in significant controversy. Significant risks have been described by withdrawing antiplatelet therapy, which included cardiovascular, cerebrovascular, and peripheral vascular thrombosis. In fact, a systematic review (69) showed a 3-fold increased risk for major adverse cardiac events and that the risk was magnified in patients with coronary stents.

Estimates suggest that greater than two-thirds of sudden cardiac events (acute coronary syndrome or sudden cardiac death) and half of the postoperative myocardial infarctions follow initiation of an arterial plaque disruption and associated thrombosis (256-261). Parallel clinical data on cerebrovascular events are not known; however, acute coronary syndrome is associated with pro-inflammatory and pro-thrombotic states, which involve elevated levels of fibrinogen, C-reactive protein, and plasminogen activator inhibitor (262). In the postoperative period, endogenous catecholamines, increased platelet adhesiveness, and decreased fibrinolysis increase the risk of an acute coronary syndrome (256,260,263).

There is evidence that discontinuation or interruption of antiplatelet therapy may result in hypercoagulability with thrombosis or bleeding (69,264-273). Studies have determined that maintaining antiplatelet therapy throughout surgery is associated with an increase in blood loss of 2.5% to 20% with aspirin and 30% to 50% with a combined therapy of aspirin and clopidogrel (273,274). Importantly, this increased blood loss has not been shown to correlate with increased mortality, except in intracranial surgery (267,273). Reports indicate an average increase of 30% in the transfusion rate, with a complication rate from red blood cell transfusion of 0.4% (275) and mortality linked directly to massive surgical blood loss in less than 3% of reported clinical data (276). However, an important recognized risk of withdrawing antiplatelet therapy has been described via a rebound effect with increased platelet adhesiveness (277-280). This phenomenon is also a consequence of the systemic inflammatory response and the acute phase reaction to surgery, which also increases platelet adhesiveness and decreases fibrinolysis (257,260,281). It should also be noted that comorbid pathologies, such as carcinoma and diabetes, can also contribute to the prothrombotic state. Death and infarction rates in acute

coronary syndrome were doubled in the setting of drug withdrawal (279). Further, during the important period of re-endothelialization after coronary stent implantation, myocardial infarction rates up to 85% have been reported (282), along with a 5-10 fold increase in the perioperative mortality. In comparison, existing literature has reported average stent thrombosis of 35% and the average mortality from stent thrombosis of 20% to 40% (257,283) without noncardiac surgery.

The questions have been raised in reference to withholding antithrombotic therapy prior to interventions and associated risks. Chassot et al (257) proposed to modify the practice of withholding patients from all antiplatelet agents 7 to 10 days prior to surgery, except when bleeding might occur in a closed cavity. Chassot et al (257) conducted an extensive review of the literature and reasoned that in the absence of prior clinical data on different antiplatelet regimens during noncardiac surgery, including low coronary risk situations, patients on antiplatelet drugs should continue therapy throughout surgery, except when bleeding might occur in a closed space. Another proposed consideration is the use of a therapeutic bridge utilizing antiplatelet drugs with shorter durations of action. In fact, other comprehensive reviews have offered variable guidance (27,31,32,34,49,54,124-129,147,149,212,251,284-299) on anticoagulation management. One systematic review (299), which evaluated 31 reports determined that for dental procedures, arthrocentesis, cataract surgery, and diagnostic endoscopy most patients can proceed without alteration to their anticoagulant therapy. Several publications also delineated a variety of other procedures, including endoscopy (300,301), peripheral vascular surgery (302), and ophthalmic surgery (303) with no evidence of increased bleeding when patients are continued on antiplatelet therapy throughout the perioperative period. Gerstein et al (304) also showed that for many other procedures, minimal perioperative bleeding risk occurred when aspirin was continued, contrasted with concomitant thromboembolic risks associated with aspirin withdrawal.

The risk of thromboembolic events may be a more significant clinical feature, versus the risk of epidural hematoma. A majority of regional anesthesia and interventional pain management literature has focused on the risks of epidural hematoma, paying only scant attention to thromboembolic events. However, wider evidence in the literature defines multiple guidelines from other specialties that described continuation of

antiplatelet therapy during surgical interventions, including peripheral vascular surgery, with some high-risk exceptions (267-277,282,299-303,305).

These data suggest the traditional attitude of discontinuing anti-clotting medications 10 days before interventions poses considerable clinical risk (268-273,277,282). Multiple prior guidelines have recommended discontinuation of anti-thrombosis therapy and this has shaped general practice to discontinue these drugs (40,174,251). Thus, despite absence of evidence of significant bleeding risk during interventional techniques in patients with antithrombotic therapy (37,251,284,285), they are routinely discontinued. Importantly, neither continuation nor discontinuation of antithrombotic therapy comes without risk (199,200,212,251,306-311). While most available clinical case reports focus on regional anesthesia for surgical procedures, additional evidence have been provided in the literature relating to epidural hematoma in patients undergoing interventional techniques for chronic pain with or without antithrombotic therapy - continued or discontinued (31,32,37,38,113-116,182,183,185-191,193,195,196,199,200,212,251,284,285,306-312).

In fact, the evidence based on this analysis for discontinuing antithrombotic agents is very weak. Evidence is stronger for the continuation for antithrombotic therapy since the majority of epidural hematomas have occured in patients either not on antithrombotic agents or in patients whose therapy had been appropriately discontinued.

6.2 Anatomical Considerations

Multiple factors herein described, including anatomy of the epidural space, spinal anatomy, coexisting pathology, coagulation status, needle placement, and injectate can affect the development of hematomas. A meta-analysis of over 600 spinal hematomas including both spontaneous and iatrogenic cases, revealed anticoagulation to be the second most common underlying etiology, closely following idiopathic occurrence without identifiable cause (313). In this meta-analysis, spinal and epidural anesthetic procedures in combination with anticoagulation represented the fifth highest cause of epidural hematomas. However, among this group of patients, spinal hematomas were most commonly encountered in those receiving continuous and not single-shot epidural anesthesia (314,315). As shown in Table 10, Lagerkranser (52) has shown patient-related risk factors in their study of 166 case reports of spinal hematoma from 1994 through 2015. Spinal stenosis was

the most common diagnosis with ankylosing spondylitis taking the third place in relation to anatomic factors.

The evidence from the present analysis also shows spinal stenosis as a major risk factor associated with technical factors of bleeding and multiple attempts.

6.3 Drug-Related Risk Factors

Drug-related factors are considered to be the most common risk factors. Continuation or discontinuation of antiplatelets and antithrombotic drugs was reported in the majority of the patients in this assessment (63%). In addition, earlier studies also showed these as the most common risk factors. Lagerkranser (52) described drug-related factors as the highest with frequencies as shown in Table 11. He reported intake of anti-hemostatic drugs in 63% of the patients, whereas more than one anti-hemostatic drug was implicated in 47 of 166 cases.

6.4 Bridge Therapy

Those undergoing chronic anticoagulation often require interruptions due to a procedure or other medical issue for various durations. The decision to bridge with heparin or other anticoagulants is a common clinical problem for those undergoing oral anticoagulation, especially in the case of interventional anesthesia. The evidence to inform decision making is limited, making current guidelines equivocal and imprecise. New randomized controlled studies have shown significant perioperative or periprocedural bleeding rates without reduction in thromboembolism when bridging is completed (314-317). Additional studies such as PERIOP2 (318) outlining guidelines for high risk individuals in need for bridging have made it somewhat clearer that only those with significant thromboembolism risk benefit from bridge therapy. Bridging anticoagulation may be vital for patients with a high risk for thromboembolism, but for most patients it produces excessive bleeding which leads to longer hospital stays and significant comorbidities without any difference in prevention of thromboembolism (319).

6.5 Reversal of Antithrombotic and Anticoagulant Agents

While immediate reversal for interventional pain procedures is rare, anticoagulation reversal and treatment options should be discussed in relation to major bleeding. Options include protamine, phytonadione (Vitamin K), hemodialysis, oral-activated charcoal, antifibrinolytic agents including tranexamic acid,

desmopressin, blood products including packed red blood cells (PRBCs) and platelets, prothrombin complex concentrates (PCCs), and specific reversal agents. For example, Idarucizumab is the only FDA-approved reversal agent for direct oral anticoagulants, specifically dabigatran. Andexanet alfa does the same for both oral and injectable factor Xa inhibitors. Prothrombin complex concentrates have been developed to contain highly concentrated coagulation factors along with antithrombotic agents. They are quite successful in the reversal of vitamin K antagonists (VKAs) and are considered first line treatment in patients with major bleeding on VKAs. Fresh Frozen Plasma traditionally has been used for VKAs as it contains all members of the coagulation cascade. It is not as beneficial for direct oral anticoagulants as these medications directly target and inactivate specific factors. The amount of FFP needed is estimated to be nearly two liters in order to appropriately establish previous concentrations of the respective factor (320).

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Conflicts of Interest

- Dr. Kaye is a speaker for Depomed, Inc. and Merck. Dr. Manchikanti has provided limited consulting services to Semnur Pharmaceuticals, Incorporated, which is developing nonparticulate steroids.
- Dr. Grider is a consultant for Medtronic and Abbott and is an Intralink Spine medical advisory board shareholder.
- Dr. Harned receives payment for development of educational presentations from Medtronic and honorarium frm St. Jude Medical.
- Dr. Benyamin is a paid consultant for Medtronic Inc., Vertiflex Inc., Vertos Medical Inc.
- Dr. Simopoulos is a consultant for Nevro Corp, St. Jude Medical, and Boston Scientific.
- Dr. Urman reports grants and personal fees from Merck, grants from Mallinckrodt, grants from Medtronic, outside the submitted work.
- Dr. Deer is a consultant for Axonics, Bioness, Abbott, Nevro, Jazz Pharm, SpineThera, Saluda, Medtronic, Vertos, Flowonix, and Ethos; minor stock/stock options with Axonics, Bioness, SpineThera, Saluda, Vertos, and Ethos; travel/accommodations expenses covered or reimburse by Abbott and Axonics; and has funded research with Saluda, Nevro, Mainstay, Abbott, Medtronic, Bioness, and JazzPharm.
 - Dr. Beall receives consulting fees from MII.
 - Dr. Hirsch is a consultant for Medtronic.

APPENDIX

Appendix Table 1

 $Appendix\ Table\ 2$

 $Appendix\ Table\ 3$

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