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ORIGINAL ARTICLE

#### **Observational Study**

## Therapeutic anticoagulation for splanchnic vein thrombosis in acute pancreatitis: A national survey and case-vignette study

Noor J Sissingh, Jesse V Groen, Hester C Timmerhuis, Marc G Besselink, Bas Boekestijn, Thomas L Bollen, Bert A Bonsing, Frederikus A Klok, Hjalmar C van Santvoort, Robert C Verdonk, Casper H J van Eijck, Jeanin E van Hooft, Jan Sven D Mieog

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#### **Abstract**

#### **BACKGROUND**

Splanchnic vein thrombosis (SVT) is a major complication of moderate and severe acute pancreatitis. There is no consensus on whether therapeutic anticoagulation should be started in patients with acute pancreatitis and SVT.

To gain insight into current opinions and clinical decision making of pancreatologists regarding SVT in acute pancreatitis.

#### **METHODS**

A total of 139 pancreatologists of the Dutch Pancreatitis Study Group and Dutch Pancreatic Cancer Group were approached to complete an online survey and case vignette survey. The threshold to assume group agreement was set at 75%.

#### RESULTS

The response rate was 67% (n = 93). Seventy-one pancreatologists (77%) regularly prescribed therapeutic anticoagulation in case of SVT, and 12 pancreatologists (13%) for narrowing of splanchnic vein lumen. The most common reason to treat SVT was to avoid complications (87%). Acute thrombosis was the most important factor to prescribe therapeutic anticoagulation (90%). Portal vein thrombosis was chosen as the most preferred location to initiate therapeutic anticoagulation (76%) and splenic vein thrombosis as the least preferred location (86%). The preferred initial agent was low molecular weight heparin (LMWH; 87%). In the case vignettes, therapeutic anticoagulation was prescribed for acute portal vein thrombosis, with or without suspected infected necrosis (82% and 90%), and thrombus progression (88%). Agreement was lacking regarding the selection and duration of long-term anticoagulation, the indication for thrombophilia testing and upper endoscopy, and about whether risk of bleeding is a major barrier for therapeutic anticoagulation.

#### **CONCLUSION**

In this national survey, the pancreatologists seemed to agree on the use of therapeutic anticoagulation, using LMWH in the acute phase, for acute portal thrombosis and in the case of thrombus progression, irrespective of the presence of infected necrosis.

Key Words: Acute pancreatitis; Splanchnic vein thrombosis; Therapeutic anticoagulation; Bleeding; Recanalization; Outcomes

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Core Tip: Splanchnic vein thrombosis is a relatively common complication of moderate and severe acute pancreatitis, but there is still much debate about its treatment with therapeutic anticoagulation. This national survey and case vignette study among 93 pancreatologists demonstrates that the majority prescribe therapeutic anticoagulation for acute portal vein thrombosis and thrombus progression in patients with or without infected necrosis, despite the absence of evidence supporting its use. Whether this collective opinion is accurate needs to be confirmed in future (preferably prospective) studies.

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#### INTRODUCTION

Acute pancreatitis is an inflammatory disorder of the pancreas and is self-limiting in the majority of patients [1,2]. However, approximately 20% of patients develop a moderate or severe disease course, with (peri) pancreatic necrosis and collections[3,4]. Due to the combination of local inflammation and



mechanical compression, these complications may cause thrombus formation in the splanchnic circulation, including the portal, splenic and superior mesenteric vein[5,6]. The reported estimates on the incidence of splanchnic vein thrombosis (SVT) in acute pancreatitis range from 17% to 23%, and are even higher in complicated acute pancreatitis [7,8]. The clinical presentation of SVT varies between an asymptomatic thrombus to potential lethal complications, such as portal or left side hypertensive bleeding and small bowel ischemia[9-11]. For this reason, early treatment with therapeutic anticoagulation is recommended in patients with acute SVT[12-14]. However, consistent evidence to drive this decision in acute pancreatitis patients does not exist[15-18]. In fact, a recent meta-analysis from our study group showed that 53% of acute pancreatitis patients do not receive therapeutic anticoagulation [15]. This proportion of untreated patients is substantially higher than previously reported in other SVT populations[19], probably because of the fear of serious bleeding. Variation in clinical practice also became apparent in this meta-analysis[15], as anticoagulation use and the type of agent used were very heterogeneous between studies. Therefore, the aim of this survey was to gain more insight into current opinions of pancreatologists on anticoagulation therapy for SVT following acute pancreatitis.

#### MATERIALS AND METHODS

We conducted an online national survey and case vignette study among members of the Dutch Pancreatitis Study Group (DPSG) and the Dutch Pancreatic Cancer Group (DPCG). Members were excluded if they were not primary care-takers in the treatment of patients with AP (e.g., radiologists, oncologists, basic scientists). The survey was built in Research Electronic Data Capture, and invitations to participate were sent by e-mail in November 2021, followed by four weekly reminders. Additionally, the survey was promoted through newsletters and during annual study group meetings of the DPSG and DPCG.

#### Survey design

The survey was developed by a multidisciplinary team of surgeons, gastroenterologists, and radiologists, and included 3 demographical questions, 17 general questions and 3 case vignettes (Supplementary material). Demographic information included the responders' specialty, type of hospital and working experience. The general questions focused on treatment of SVT and potential factors that may influence the responders' decision. The case-vignettes addressed the preferred treatment strategy in different clinical cases at different time points. All cases however, concerned a 50-year-old male patient with acute alcoholic necrotising pancreatitis, and can be summarized as follows.

Case vignette 1: A patient visited the emergency department, 5 d after onset of abdominal pain. Contrast-enhanced CT (CECT) showed necrotising pancreatitis with acute necrotic collection in the head of the pancreas (Figure 1A) and: A1: Luminal narrowing of the portal vein without the presence of collateral circulation (Figure 1B); A2: Intraluminal filling defect in the portal vein without the presence of collateral circulation; A3: Intraluminal filling defect in the portal vein without the presence of collateral circulation + a pseudoaneurysm in the proximal splenic artery (Figure 1C).

Case vignette 2: A patient admitted to the ward, 14 d after onset of abdominal pain. The patient showed signs of clinical detoriation with fever and rising inflammatory parameters. The CECT showed almost fully encapsulated pancreatic necrosis without gas configurations (Figure 1D) and a new intraluminal filling defect in the portal vein without the presence of collateral circulation (Figure 1E). The diagnosis of suspected infected pancreatic necrosis was made.

Case vignette 3: A homeless patient visited the emergency department, 30 d after onset of vague abdominal pain. CECT showed necrotising pancreatitis and: CA: Intraluminal filling defect in the portal vein with the presence of collateral circulation; CB: Thrombus progression and expansion of the collateral circulation (Figure 1F).

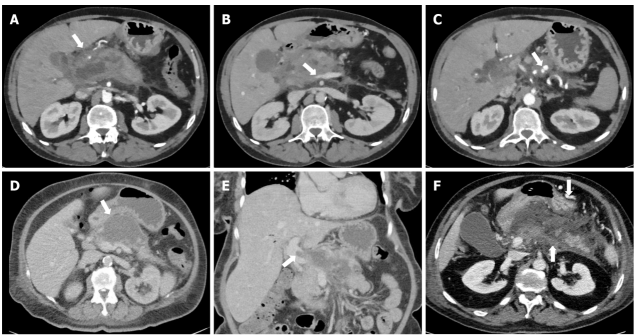
The threshold to assume group agreement was set at 75%. If a question ranged from always, usually, sometimes and never, agreement was defined when 75% of the pancreatologists rated it as always or usually (regularly), or sometimes and never (infrequently).

#### **Definitions**

SVT was predefined as an actual intraluminal filling defect on imaging of one or more of the splanchnic veins. The chronicity was divided into (sub)acute thrombosis or chronic thrombosis (with concomitant collaterals), anatomical location into portal, splenic and/or superior mesenteric vein, degree into a total or partial occlusion and extent into an isolated thrombus or a thrombus in several venous segments. Thrombus progression was defined as progression into other splanchnic vein(s), into total occlusion, or

#### Statistical analysis

Descriptive data are presented as counts with proportions for categorical data. All analyses were performed using IBM SPSS (20).



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Figure 1 Imaging findings of case vignette. A: Acute necrotic collection in the head of the pancreas in case vignette 1; B: Luminal narrowing of the portal vein without the presence of collateral circulation in case vignette 1; C: Pseudoaneurysm in the proximal splenic artery in case vignette 1; D: Almost fully encapsulated pancreatic necrosis without gas configurations in case vignette 2; E: Luminal filling defect in the portal vein without the presence of collateral circulation in case vignette 2; F: Extension of the thrombus to the splenic vein (arrow pointing upwards) and expansion of the collateral pathway in the gastroepiploic veins along the great curvature of the stomach (arrow pointing downwards) in case vignette 3.

#### RESULTS

A total of 93 of the 139 invited pancreatologists (67%) responded and participated in this survey and case vignette study; 67 gastroenterologists (72%), 25 surgeons (27%) and 1 intensivist (1%). The majority worked in a non-academic centre (70%) and had more than 10 years of experience in treating AP patients (60%). Demographic characteristics are presented in Table 1.

#### Indications for and details of treatment with therapeutic anticoagulation

Agreement was reached on whether therapeutic anticoagulation should be prescribed for SVT and luminal narrowing of one or more of the splanchnic veins in acute pancreatitis patients. For SVT, therapeutic anticoagulation was regularly prescribed by 71 (76%) and infrequently by 22 (24%) pancreatologists. In case of luminal narrowing, therapeutic anticoagulation was only regularly prescribed by 12 (13%) pancreatologists. Avoiding complications, such as portal hypertension and bowel ischemia, was the main reason for 81 pancreatologists (87%) to start therapeutic anticoagulation. Screening for an underlying prothrombotic disorder in patients diagnosed with SVT was regularly performed by 14 (15%) pancreatologists, only in patients with a history of one (or more) thrombotic events by 40 (43%), and infrequently by 39 (42%) pancreatologists. There was agreement on the preferred initial type of therapeutic anticoagulation for SVT 81 pancreatologists (87%) preferred subcutaneous low-molecularweight heparin (LMWH)], but not on the preferred follow-up type. Imaging after the index admission was chosen as follow-up strategy by 79 pancreatologists (85%). Thirteen pancreatologists (13%) indicated that they usually stop anticoagulant therapy in case of achieved radiological recanalization, 35 (38%) after a period of 3 mo, 42 (45%) after 6 mo, and 3 (3%) after 12 mo. All details are provided in

#### Determinants of prescribing therapeutic anticoagulation

Seventy-eight pancreatologists (84%) have chosen the time course of thrombosis as the most important factor supporting anticoagulant therapy; 84 pancreatologists (90%) prescribe therapeutic anticoagulation in case of a (sub)acute thrombosis vs 9 (10%) for both (sub)acute and chronic thrombosis. Moreover, 70 pancreatologists (76%) have chosen portal vein thrombosis as the most preferred location to initiate therapeutic anticoagulation, whereas splenic vein thrombosis was chosen as least preferred location by 80 pancreatologists (86%). The majority of pancreatologists (85%) treat both total and partial occlusive thrombosis. There was no agreement whether the risk of different types of bleeding should be considered as a major barrier to prescribe therapeutic anticoagulation. The need for invasive interventions for local complications of acute pancreatitis influenced the decision whether or not to

Table 1 Details of respondents, n (%)	
Demographics	n = 93
Specialty	
Surgeon	25 (27)
Gastroenterologist	67 (72)
Intensivist	1 (1)
Type of hospital	
Academic	28 (30)
Non-academic, teaching hospital	60 (65)
Non-academic, non-teaching hospital	5 (5)
Experience in treating patients with acute pancreatitis	
0-5 years	10 (11)
5-10 years	27 (29)
10-15 years	17 (18)
15-20 years	23 (25)
> 20 years	16 (17)

initiate anticoagulation therapy in about half of pancreatologists (52%). All details are outlined in Table 3.

#### Statements on prognosis

An association between the presence of SVT and worse clinical outcomes in patients with acute pancreatitis was assumed by 67 pancreatologists (72%) (Figure 2). Moreover, the vast majority (88%) agreed that therapeutic anticoagulation for splanchnic vein thrombosis improves clinical outcomes in these patients. Insufficient evidence was the most frequently quoted reason among pancreatologists who disagreed with this second statement.

#### Case-vignettes

The results of the case vignettes are summarized in Figure 3. In the first case vignette (patient 1, day 5 of acute necrotising pancreatitis), 11 pancreatologists (12%) would prescribe a therapeutic dose anticoagulation if luminal narrowing without collateral circulation was detected in the portal vein. Of the 82 pancreatologists (88%) who opted for no therapeutic dose anticoagulation, 73 (89%) would change treatment strategy in case an actual filling defect in the portal vein was detected. In total, 84 pancreatologists (90%) would prescribe therapeutic dose anticoagulation to this patient with an actual portal vein thrombosis without collateral circulation. If a pseudoaneurysm was concomitantly present, 43 of those 84 pancreatologists (51%) who favoured a therapeutic dose anticoagulation would switch to a prophylactic dose anticoagulation (n = 28, 65%) or no anticoagulation at all (n = 15, 35%), leaving 41 pancreatologists (44%) in the therapeutic anticoagulation group.

In the second case vignette (patient 2, day 14 of suspected infected necrotising pancreatitis), 77 pancreatologists (82%) would prescribe therapeutic dose anticoagulation if a portal vein thrombosis without collateral circulation was detected. The presence of (suspected) infected pancreatic necrosis influenced the choice of anticoagulation agent in 49 pancreatologists (52%). Almost all of these pancreatologists pointed out that once infected pancreatic necrosis is suspected, they would choose an agent with a short half-life because of the potentially need of invasive intervention.

In the third case vignette (patient 3, day 30 of acute necrotising pancreatitis), 44 pancreatologists (47%) would prescribe a therapeutic dose anticoagulation if a portal vein thrombosis with collateral circulation was detected. Of these 44 pancreatologists, 19 (43%) would perform upper endoscopy to screen for and-if present-treat oesophageal varices before starting anticoagulation therapy. In case of thrombus progression (extension of the thrombus to the splenic vein and expansion of the collateral pathway), 11 pancreatologists (12%) would stay conservative (i.e., no therapeutic dose of anticoagulation), 82 (88%) would start or continue a therapeutic dose anticoagulation and none would proceed to an intervention.

#### Table 2 Survey results: Indication for and details of treatment with the rapeutic anticoagulation, n (%)

Deyrou prescribe therapeutic AC in case of detected thrombosis in one (or more) of the splanchnic veins?  Always 2625 Usually 2625 Nover 2100 Doyou prescribe therapeutic AC in case of detected furnial narrowing of one (or more) of the splanchnic veins?  Always 2610 Doyou prescribe therapeutic AC in case of detected furnial narrowing of one (or more) of the splanchnic veins?  Always 2610 Doyou Scare times devenues anatomy 2610 Doyou Scare times devenues anatomy 2610 Doyou Scare for rearrange of SVT Doyou Treamyou Color Treamyou 2610 Doyou Scare for an underlying producenholic disorder?  Always 2610 Doyou Scare for an underlying producenholic disorder?  Always 2610 Doyou Scare for an underlying producenholic disorder?  Always 2610 Doyou Scare for an underlying producenholic disorder?  Always 2610 Doyou patients with a history of one (or more) thrombosic events 2610 Doyou Scare for an underlying producenholic events 261	Item	Total (n = 93)
Usually         48 (22)           Semetimes         2 (23)           Never         1 (20)           De you prescribe therapeutic AC in case of detected luminal narrowing of one (or more) of the splanchnic veins?         150           Do you prescribe therapeutic AC in case of detected luminal narrowing of one (or more) of the splanchnic veins?         3 (5)           Locally         9 (10)           Sometimes         29 (31)           Never         5 (56)           In service veesel reconalization         5 (25)           To evoid complications         81 (87)           To prevent formation of altered venous anatomy         31 (33)           To prevent formation of altered venous anatomy         30 (32)           Other reason'         27 (29)           Use prevent formation of altered venous anatomy         30 (32)           Other reason'         10 (12)           Do you screen for an underlying prothrombotic disorder?         2 (2)           Always         2 (3)           Suisually         1 (13)           Sometimes         2 (27)           Only in patients with a history of one (or more) thrombotic events         4 (3)           Nive         1 (13)           Unfunctionated heparin intravenous         4 (4)           Unfunctionated	Do you prescribe therapeutic AC in case of detected thrombosis in one (or more) of the splanchnic veins?	
Sementimes         21 (23)           Never         1 (1)           Do you prescribe therapeutic AC in case of detected luminal narrowing of one (or more) of the splanchuic veiner?         Very Control of the splanchuic veiner?           Always         9 (10)           Sometimes         29 (31)           Never         32 (36)           Main reason() to start therapeutic AC (multiple answers were possible)         Very Control of Control of Section of allered venous anatomy         31 (33)           To aveid connalization         36 (87)           To aveid connalizations         31 (33)           To prevent formation of allered venous anatomy         31 (33)           To prevent recurrence of SVT         27 (29)           To prevent network venous thromboenbellsm         30 (32)           Other reason of         10 (10)           Do you servers for an underlying prothombetic disorder?         2 (2)           Always         2 (2)           Usually         12 (33)           Sometimes         2 (2)           Nover         14 (4)           (Low an placetians with a history of one (or more) thrombotic events         40 (32)           Never         4 (4)           (Low molecular weight) heparin subcutancous         81 (87)           (Low molecular weight) heparin sub	Always	23 (25)
Newer	Usually	48 (52)
Do you prescribe therapeutic AC in case of detected furninal narrowing of one (or more) of the splanchnic veins?         3 (3)           Always         3 (3)           Usually         9 (10)           Sometimes         29 (31)           Never         3 (56)           Main reason(s) to start therapeutic AC (multiple answers were possible)         3 (56)           In avoid complications         52 (56)           In avoid complications         31 (33)           In prevent formation of altered venous anatomy         31 (33)           In prevent recurrence of SVT         27 (29)           Other reason¹         10 (1)           Do you screen for an underlying prothrombetic disorder?         2 (2)           Always         2 (2)           Usually         2 (2)           Usually         2 (2)           Usually         2 (2)           Only in patients with a history of one (or more) thrombotic events         40 (45)           Never         40 (45)           Which initially type of therapeutic AC do you prefer?         4 (4)           (Low molecular weight) beparin subcutaneous         81 (87)           Unfractionated heparin intravenous         9 (10)           Uniforactionated heparin intravenous         0           Uniforactionated heparin i	Sometimes	21 (23)
Abways	Never	1 (1)
Samely   9 (10)   Sometimes   29 (31)   Newer   25 (26)   Main reason(e) to start therapeutic AC (multiple answers were possible)   To achieve vessel reconalization   52 (56)   To avoid complications   81 (87)   To prevent formation of altered venous anatomy   31 (33)   To prevent recurrence of SVT   27 (29)   To prevent another venous thromboembolism   30 (32)   Other rossen   10 (1)   Do you screen for an underlying prothrombotic disorder?   Always   2 (2)   Usually   3 (2)   Only in patients with a history of one (or more) thrombotic events   40 (3)   Never   40 (4)   Direct oral anticoagulation   4 (4)   Direct oral anticoagulation   4 (4)   Direct oral anticoagulation   4 (4)   Placel aggregation inhibitor   4 (4)   Urokinsocy/recombinant tissue plasminegen activator   4 (4)   Urokinsocy/recombinant tissue plasminegen activator   9 (10)   Urokinsocy/recombinant tissue plasminegen activator   9 (10)   Urifractionated heparin intravenous   9 (10)   Urifractionated apgregation inhibitor   20 (3)   Pladel aggregation inhibitor   20 (3)   Urifractionated speparin intravenous   9 (10)   Urokinsocy/recombinant tissue plasminegen activator   9 (20)   Urokinsocy/recombinant	Do you prescribe therapeutic AC in case of detected luminal narrowing of one (or more) of the splanchnic veins?	
Sometimes         29 (31)           Never         52 (56)           Main roson(s) to start therapeutic AC (multiple answers were possible)         52 (56)           To avoid complications         81 (87)           To avoid complications         81 (87)           To prevent formation of altered venous anatomy         31 (33)           To prevent another venous thrombocembolism         30 (32)           Other reason¹         1 (1)           Do you screen for an underlying prothrombotic disorder?         40 (32)           Always         2 (2)           Usually         12 (13)           Sometimes         25 (27)           Only in patients with a history of one (or more) thrombotic events         40 (43)           Never         14 (15)           Which initial type of therapeutic AC do you prefer?         44 (4)           Unfractionated heparin intravenous         4 (8)           Unfractionated heparin initravenous         4 (8)           Vitamin K antagoeist         4 (9)           Platelet aggregation inhibitor         0           Unfractionated heparin initravenous         9 (10)           Unkinses/recombinant tissue plasminogen activator         9 (10)           Unfractionated heparin initravenous         9 (10)           Uritarin K an	Always	3 (3)
Never         \$2 (50)           Main reason(s) to start therapeutic AC (multiple answers were possible)         \$2 (50)           To achieve vessel recanalization         \$2 (50)           To achieve vessel recanalization         \$1 (87)           To prevent formation of altered venous anatomy         \$1 (30)           To prevent frecurrence of SVT         \$27 (29)           To prevent another venous thromboembolism         \$0 (32)           Other reason <sup>1</sup> \$1 (1)           Do you screen for an underlying prothrombotic disorder?         \$2 (2)           Usually         \$2 (2)           Usually         \$2 (2)           Sometimes         \$2 (27)           Only in patients with a history of one (or more) thrombotic events         \$1 (15)           Never         \$1 (15)           (Low molecular weight) heparin subcutaneous         \$1 (87)           Unfractionated heparin intravenous         \$1 (8)           Unificationated heparin intravenous         \$1 (8)           Vitamin K antagonist         \$1 (8)           Unokinase/recombinant tissue plasminogen activator         \$0           Unificationated heparin intravenous         \$0           Unificationated heparin intravenous         \$0           Unificationated appreadation         \$3 (57)	Usually	9 (10)
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To achieve vessel recanalization         52 (56)           To avoid complications         81 (87)           To prevent formation of altered venous anatomy         31 (33)           To prevent recurrence of SVT         27 (29)           To prevent another venous thromboembolism         30 (32)           Other reason¹         1 (1)           Do you screen for an underlying prothrombotic disorder?         2           Always         2 (2)           Usually         25 (27)           Only in patients with a history of one (or more) thrombotic events         40 (43)           Never         14 (15)           Which initial type of therapeutic AC do you prefer?         4 (4)           Unfractionated heparin intravenous         4 (4)           Unfractionated heparin intravenous         4 (4)           Picaled aggregation inhibitor         1 (1)           Urokinase/ recombinant tissue plasminogen activator         9 (10)           Which follow-up type of therapeutic AC do you prefer?         (Low molecular weight) heparin subcutaneous         9 (10)           Unfractionated heparin intravenous         9 (10)           Unfractionated heparin intravenous         9 (20)           Unfractionated heparin intravenous         9 (20)           Direct oral anticoagulation         3 (37)	Never	52 (56)
To avoid complications         \$1 (87)           To prevent formation of altered venous anatomy         31 (33)           To prevent recurrence of SVT         27 (29)           To prevent another venous thromboembolism         30 (32)           Other reason¹         1 (1)           Do you screen for an underlying prothrombotic disorder?         ***           Always         2 (2)           Usually         12 (33)           Sometimes         25 (27)           Only in patients with a history of one (or more) thrombotic events         40 (43)           Never         40 (43)           Which initial type of therapeutic AC do you prefer?         ***           (Low molecular weight) heparin subcutaneous         4 (4)           Unfractionated heparin intravenous         4 (4)           Plated aggregation inhibitor         1 (1)           Urokinase/recombinant tissue plasminogen activator         9 (10)           Which follow-up type of therapeutic AC do you prefer?         **           (Low molecular weight) heparin subcutaneous         9 (10)           Unfractionated heparin intravenous         9 (10)           Urokinase/recombinant tissue plasminogen activator         9 (20)           Unfractionated heparin intravenous         0           Direct oral anticoagulation	Main reason(s) to start therapeutic AC (multiple answers were possible)	
To prevent formation of altered venous anatomy         31 (3)           To prevent another venous thromboembolism         30 (32)           Other reason¹         1 (1)           Do you screen for an underlying prothrombotic disorder?	To achieve vessel recanalization	52 (56)
To prevent another venous thromboembolism         30 (32)           To prevent another venous thromboembolism         30 (32)           Other reason¹         1 (1)           Do you screen for an underlying prothrombotic disorder?         ***           Always         2 (2)           Usually         12 (13)           Sometimes         25 (27)           Only in patients with a history of one (or more) thrombotic events         40 (43)           Never         44 (15)           Which initial type of therapeutic AC do you prefer?         ***           (Low molecular weight) heparin subcutaneous         81 (87)           Unfractionated heparin intravenous         4 (4)           Direct oral anticoagulation         3 (3)           Vitamin K antagonist         4 (4)           Platelet aggregation inhibitor         1 (1)           Urokinase/recombinant tissue plasminogen activator         9 (10)           Unfractionated heparin intravenous         9 (10)           Unfractionated heparin intravenous         9 (3)           Unifractionated heparin intravenous         9 (3)           Unifractionated heparin intravenous         9 (3)           Unifractionated heparin intravenous         9 (3)           Vitamin K antagonist         2 (2)	To avoid complications	81 (87)
To prevent another venous thromboembolism Other reason¹ 1(1)  Do you screen for an underlying prothrombotic disorder?  Always 2(2) Usually 12 (13) Sometimes 25 (27) Only in patients with a history of one (or more) thrombotic events Never 14 (15)  Which initial type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous 4 (4) Direct oral anticoagulation Vitamin K antagonist 4 (4) Platelet aggregation inhibitor Unfractionated heparin intravenous 0 Which follow-up type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous 0 Which follow-up type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous 0 Unfractionated heparin intravenous 0 Unifractionated heparin intravenous 0 Unifractionated heparin intravenous 0 Unfractionated heparin intravenous 0 Direct oral anticoagulation 25 (37) Vitamin K antagonist 29 (31) Platelet aggregation inhibitor 20 (2) Unokinase/recombinant tissue plasminogen activator 0 Direct oral anticoagulation 15 (37) Vitamin K antagonist 20 (21) Unokinase/recombinant tissue plasminogen activator 0 Do you generally follow-up SVT after index admission?  Yes, clinically only 5 (5) Fys, with imaging 5 (5) Fys, with imaging	To prevent formation of altered venous anatomy	31 (33)
Other reason¹         1 (1)           Do you screen for an underlying prothrombotic disorder?         2 (2)           Always         12 (13)           Sometimes         25 (27)           Only in patients with a history of one (or more) thrombotic events         40 (43)           Never         14 (15)           Which initial type of therapeutic AC do you prefer?         81 (87)           Unifractionated heparin intravenous         4 (4)           Direct oral anticoagulation         3 (3)           Vitamin K antagonist         4 (4)           Platelet aggregation inhibitor         1 (1)           Urokinase/recombinant tissue plasminogen activator         0           Which follow-up type of therapeutic AC do you prefer?         (Low molecular weight) heparin subcutaneous         9 (10)           Unfractionated heparin intravenous         9 (10)         0           Unifractionated heparin intravenous         2 (3)           Vitamin K antagonist         29 (31)           Platelet aggregation inhibitor         2 (2)           Urokinase/recombinant tissue plasminogen activator         0           Do you generally follow-up SVT after index admission?         2 (2)           Yes, clinically only         5 (5)           Yes, with imaging         79 (85)	To prevent recurrence of SVT	27 (29)
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Always       2 (2)         Usually       12 (13)         Sometimes       25 (27)         Only in patients with a history of one (or more) thrombotic events       40 (43)         Never       14 (15)         Which initial type of therapeutic AC do you prefer?       ***         (Low molecular weight) heparin subcutaneous       \$1 (87)         Unfractionated heparin intravenous       4 (4)         Direct oral anticoagulation       3 (3)         Vitamin K antagonist       4 (4)         Platelet aggregation inhibitor       1 (1)         Urokinase/recombinant tissue plasminogen activator       9 (10)         Which follow-up type of therapeutic AC do you prefer?       9 (10)         (Low molecular weight) heparin subcutaneous       9 (10)         Unfractionated heparin intravenous       9 (10)         Direct oral anticoagulation       53 (57)         Vitamin K antagonist       29 (31)         Platelet aggregation inhibitor       2 (2)         Urokinase/recombinant tissue plasminogen activator       0         Do you generally follow-up SVT after index admission?       5 (5)         Yes, clinically only       5 (5)         Yes, with imaging       79 (85)	Other reason <sup>1</sup>	1 (1)
Usually12 (13)Sometimes25 (27)Only in patients with a history of one (or more) thrombotic events40 (43)Never14 (15)Which initial type of therapeutic AC do you prefer?81 (87)(Low molecular weight) heparin subcutaneous81 (87)Unfractionated heparin intravenous4 (4)Direct oral anticoagulation3 (3)Vitamin K antagonist4 (4)Platelet aggregation inhibitor1 (1)Urokinase/ recombinant tissue plasminogen activator0Which follow-up type of therapeutic AC do you prefer?9 (10)(Low molecular weight) heparin subcutaneous9 (10)Unfractionated heparin intravenous5 (35)Direct oral anticoagulation53 (57)Vitamin K antagonist29 (31)Platelet aggregation inhibitor2 (2)Urokinase/ recombinant tissue plasminogen activator20Do you generally follow-up SVT after index admission?5 (5)Yes, clinically only5 (5)Yes, with imaging79 (85)	Do you screen for an underlying prothrombotic disorder?	
Sometimes 25 (27) Only in patients with a history of one (or more) thrombotic events 40 (43) Never 14 (15) Which initial type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous 81 (87) Unfractionated heparin intravenous 44 (4) Direct oral anticoagulation 3 (3) Vitamin K antagonist 4 (4) Platelet aggregation inhibitor 1 (1) Urokinase/recombinant tissue plasminogen activator 0 Which follow-up type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous 9 (10) Unfractionated heparin intravenous 53 (57) Vitamin K antagonist 22 (2) Urokinase/recombinant tissue plasminogen activator 22 (2) Urokinase/recombinant tissue plasminogen activator 55 (5) Dy you generally follow-up SVT after index admission?  Yes, clinically only 5 (8)	Always	2 (2)
Never 14 (15)  Never 14 (15)  Which initial type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous 81 (87)  Unfractionated heparin intravenous 4 (4)  Direct oral anticoagulation 3 (3)  Vitamin K antagonist 4 (4)  Platelet aggregation inhibitor 1 (1)  Urokinase/recombinant tissue plasminogen activator 0 0  Which follow-up type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous 9 (10)  Unfractionated heparin intravenous 0 9 (10)  Unfractionated heparin intravenous 9 (10)  Unfractionated heparin intravenous 9 (10)  Unfractionated heparin intravenous 9 (35)  Vitamin K antagonist 29 (31)  Platelet aggregation inhibitor 20 (2)  Urokinase/recombinant tissue plasminogen activator 20 (3)  Vitamin K antagonist 20 (3)  Platelet aggregation inhibitor 20 (2)  Urokinase/recombinant tissue plasminogen activator 5 (5)  Do you generally follow-up SVT after index admission?  Yes, clinically only 5 (5)  Ps, with imaging 79 (85)	Usually	12 (13)
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Which initial type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous 4 (4)  Direct oral anticoagulation 3 (3)  Vitamin K antagonist 4 (4)  Platelet aggregation inhibitor 1 (1)  Urokinase/recombinant tissue plasminogen activator  Which follow-up type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous 9 (10)  Unfractionated heparin intravenous 9 (10)  Unfractionated heparin intravenous 9 (10)  Unfractionated heparin intravenous 9 (10)  Unfractionated pagregation inhibitor 2 (2)  Urokinase/recombinant tissue plasminogen activator 0  Do you generally follow-up SVT after index admission? Yes, clinically only 9 (85)	Only in patients with a history of one (or more) thrombotic events	40 (43)
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Vitamin K antagonist 4 (4)  Platelet aggregation inhibitor 1 (1)  Urokinase/recombinant tissue plasminogen activator 0  Which follow-up type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous 9 (10)  Unfractionated heparin intravenous 0  Direct oral anticoagulation 53 (57)  Vitamin K antagonist 29 (31)  Platelet aggregation inhibitor 2 (2)  Urokinase/recombinant tissue plasminogen activator 0  Do you generally follow-up SVT after index admission?  Yes, clinically only 5 (5)  Yes, with imaging 79 (85)	Unfractionated heparin intravenous	4 (4)
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Urokinase/recombinant tissue plasminogen activator  Which follow-up type of therapeutic AC do you prefer?  (Low molecular weight) heparin subcutaneous  9 (10)  Unfractionated heparin intravenous  0  Direct oral anticoagulation  53 (57)  Vitamin K antagonist  29 (31)  Platelet aggregation inhibitor  2 (2)  Urokinase/recombinant tissue plasminogen activator  0  Do you generally follow-up SVT after index admission?  Yes, clinically only  5 (5)  Yes, with imaging	Vitamin K antagonist	4 (4)
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Vitamin K antagonist  Platelet aggregation inhibitor  2 (2)  Urokinase/recombinant tissue plasminogen activator  Do you generally follow-up SVT after index admission?  Yes, clinically only  Yes, with imaging  29 (31)  2 (2)  0  79 (85)	Unfractionated heparin intravenous	0
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Urokinase/recombinant tissue plasminogen activator  Do you generally follow-up SVT after index admission?  Yes, clinically only  Yes, with imaging  79 (85)	Vitamin K antagonist	29 (31)
Do you generally follow-up SVT after index admission?  Yes, clinically only  Yes, with imaging  5 (5)  79 (85)	Platelet aggregation inhibitor	2 (2)
Yes, clinically only  Yes, with imaging  79 (85)	Urokinase/recombinant tissue plasminogen activator	0
Yes, with imaging 79 (85)	Do you generally follow-up SVT after index admission?	
	Yes, clinically only	5 (5)
No 9 (10)	Yes, with imaging	79 (85)
	No	9 (10)

After how long do you usually stop the therapeutic AC?	
In case of achieved radiological recanalization	13 (14)
3 mo	35 (38)
6 mo	42 (45)
12 mo	3 (3)
Never	0

<sup>&</sup>lt;sup>1</sup>In free text: expansion of thrombosis.

AC: Anticoagulation; SVT: Splanchnic vein thrombosis.

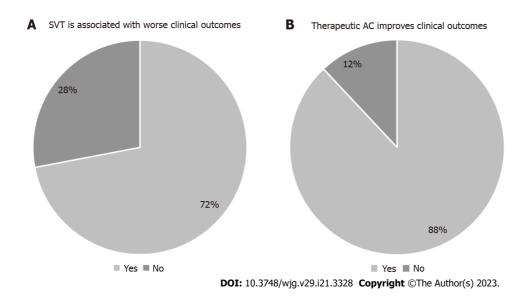


Figure 2 Statements on prognosis. A: Splanchnic vein thrombosis is associated with worse clinical outcomes; B: Therapeutic anticoagulation improves clinical outcomes. AC: Anticoagulation; SVT: Splanchnic vein thrombosis.

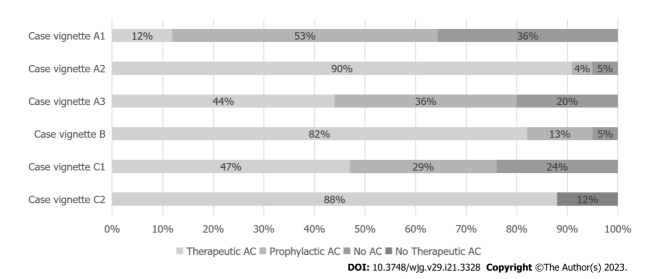


Figure 3 Case vignettes results: Choice of treatment. AC: Anticoagulation.

#### DISCUSSION

This first nationwide survey and case vignette study gives insight into the clinical scenarios in which therapeutic anticoagulation is currently used, and not used to treat or prevent splanchnic vein thrombosis in acute pancreatitis patients. In an earlier study [15], we found 7 retrospective cohort studies evaluating therapeutic anticoagulation in this patient category with conflicting results in clinical

#### Table 3 Survey results: Determinants of prescribing the rapeutic anticoagulation, n (%)

Item	Total (n = 93)
Do you consider of the thrombosis as an important factor to prescribe therapeutic AC? (multiple answers were possible)	
Age (acute or chronic)	78 (84)
Anatomical location (portal, splenic or superior mesenteric vein)	42 (45)
Degree (total or partial)	45 (48)
Extent (isolated thrombosis or thrombosis in several segments)	49 (53)
Progression (over time)	40 (43)
When do you prescribe therapeutic AC? In case of:	
(Sub)acute thrombosis	84 (90)
Chronic thrombosis	0
Both	9 (10)
Rank the anatomical location of the thrombosis from most likely to less likely to start therapeutic AC:	
Portal vein-splenic vein-superior mesenteric vein	9 (10)
Portal vein-superior mesenteric vein-splenic vein	61 (66)
Splenic vein-portal vein-superior mesenteric vein	0
Splenic vein-superior mesenteric vein-portal vein	1 (1)
Superior mesenteric vein-portal vein-splenic vein	19 (20)
Superior mesenteric vein-splenic vein-portal vein	3 (3)
When do you prescribe therapeutic AC? In case of:	
Total thrombosis	9 (10)
Partial thrombosis	5 (5)
Both	79 (85)
Do you consider the risk of as a major barrier to prescribe therapeutic AC? (multiple answers were possible)	
Bleeding in general	52 (56)
Bleeding related to portal hypertension	17 (18)
Bleeding related to pseudoaneurysm	49 (53)
Other risk <sup>1</sup>	1 (1)
Does the need for invasive interventions for local complications of acute pancreatitis influence your decision regarding AC therapy?	
Yes	48 (52)
No	45 (48)

<sup>1</sup>In free text: CVA bleeding history.

AC: Anticoagulation; SVT: Splanchnic vein thrombosis.

outcome [20-27]. These studies were of moderate quality and therefore the pancreatologist' preference and belief predominate in current decision making rather than scientific evidence.

An important finding of the current study was that more than 75% of pancreatologists regularly prescribe therapeutic anticoagulation for SVT, particularly for a thrombus that acutely developed. This is in line with recommendations from general guidelines for SVT management [12-14]. In the absence of a visualized thrombus, most pancreatologists indicated not to treat compressed veins with anticoagulation. Although wall shear stress in a compressed vessel may promote platelet activation, and subsequently thrombus formation[28], there is no data yet to question the opinion of the pancreato-

In this study, the most important reason to administer therapeutic anticoagulation was to avoid complications including bowel ischemia and portal hypertension. Bowel ischemia has been reported in up to 33% of acute pancreatitis patients treated with therapeutic anticoagulation vs 16% of untreated patients[22,24,25]. A potential explanation for this discrepancy could be that bowel ischemia was already present prior to the start of therapy, therefore being an indication for therapeutic anticoagulation rather than a consequence. In addition, the presence of varices and other collaterals have been equally reported[15], and only one of the aforementioned studies described one case of bleeding from oesophageal varices in an anticoagulated patient[24]. Again, it is likely that a perceived bleeding risk influenced the decision whether or not to prescribe therapeutic anticoagulation. This confounding by indication clearly limits the interpretation of these retrospective studies.

Achieving vessel recanalization was chosen as the second goal. A recent meta-analysis showed that the pooled rate of recanalization of SVT was similar between treated (36%) and untreated patients (31%) [15]. However, there is reason to believe that the benefit of anticoagulation therapy may alter when considering the anatomical location of the thrombosis[21,29]. Patients with portal vein or superior mesenteric vein thrombosis may have an increased risk of complications, while having lower spontaneous recanalization rates. In particular, mortality rates of patients with superior mesenteric vein thrombosis are reported up to 50% [30,31]. On the other hand, splenic vein thrombosis, which is by far the most common site of thrombosis in acute pancreatitis patients, forms a less serious concern for gastrointestinal bleeding and insufficient recanalization [8,26,32]. A selective anticoagulation policy, in which therapeutic anticoagulation was reserved for portal- and superior mesenteric vein thrombosis, was recently assessed in a retrospective study [33]. This study showed a recanalization rate of 67% in portal- and superior mesenteric vein thrombosis, which is substantially higher than previously reported [15]. In addition, a recent practice guideline from the Pancreas study group, Chinese Society of Gastroenterology, recommends a selective anticoagulation policy[34]. In this survey, portal vein thrombosis, followed by superior mesenteric vein thrombosis, was also the pancreatologists' preferred location for prescribing therapeutic anticoagulation, while splenic vein was the least preferred location.

With respect to chronic SVT, the current guidelines do not recommend therapeutic anticoagulation [10]. This is in line with the reported use in case vignette 3, with the exception of the case of the patient with thrombus progression and expansion of the collateral circulation. In this scenario, 88% of pancreatologists would treat such patient with therapeutic anticoagulation. A recent multicentre randomised controlled trial comparing daily rivaroxaban 15 mg/d to no anticoagulation in patients with noncirrhotic chronic portal vein thrombosis [35], formally challenged the guideline recommendations. This study showed that rivaroxaban, even in prophylactic dose, reduced the incidence of venous thromboembolism; therefore, this study may initiate a shift towards a more frequent use of anticoagulation in chronic SVT. On that note, primary prophylaxis of portal hypertensive bleeding should be performed, as laid out by the BAVENO IV guideline[13]. In this survey, however, the minority of pancreatologists followed this recommendation. Improvements should also be made to distinguish acute from chronic SVT. Currently, no clear definition for chronic SVT exists other than a presumed time course of more than 6 mo or the presence of multiple small collaterals around the obstructed veins [10,36], which is not useful to diagnose a nonocclusive chronic thrombosis (i.e., absence of collateral pathways). A promising invention to overcome this problem is magnetic resonance noncontrast thrombus imaging, though validation is still needed[37].

According to our survey, subcutaneous LMWH was the favoured initial type of therapeutic anticoagulation, while no agreement regarding the choice of long-term anticoagulation and its duration was found. In current guidelines, switching LMWH to a vitamin K antagonist once reaching the target range is the reported strategy for patients with SVT[12-14,38]. The use of direct oral anticoagulation (i.e., apixaban) in acute pancreatitis patients with SVT is reported in two studies and showed comparable results [21,33]. However, in the case of (suspected) infected pancreatic necrosis, LMWH seems to be preferred by more than half of the pancreatologists, due to its short half-life and reversibility. Besides, many acute pancreatitis patients have reduced caloric intake limiting the absorption of DOACS. Therefore, it seems fair to advise LWMH, especially in the acute phase. Looking at the duration of anticoagulation therapy for provoked SVT in patients with a transient risk factor, such as acute pancreatitis, the suggested duration is 3 mo to 6 mo[12-14,38]. Consistently, 38% and 45% of the pancreatologists in our survey preferred 3 mo and 6 mo treatment duration, respectively.

Based on the available literature, it remains unclear whether therapeutic anticoagulation is associated with higher rates of bleeding. An increased bleeding risk with therapeutic anticoagulation has been reported up to 33% of patients[21,25,26], but there are also studies showing lower rates of bleeding[24]. The theory for this latter finding is that therapeutic anticoagulation prevents thrombus progression, therefore reducing the portal pressure and consequently the risk of bleeding [19]. In this study, the risk of bleeding was not identified as a significant discouraging factor, as only about half of the pancreatologists considered bleeding in general and bleeding related to pseudoaneurysm as a major barrier to prescribe therapeutic anticoagulation. Also, the possible need for invasive intervention, due to suspected infected necrosis, did not significantly influence the treatment strategy. Another critical question is whether SVT influences the disease course of acute pancreatitis patients, but again this remains unanswered[15]. In this survey, the majority of pancreatologists assumed that the occurrence of SVT is associated with worse clinical outcomes, and interestingly, even more pancreatologists were convinced that the use of therapeutic anticoagulation leads to improved patient outcomes.

A strength of this study is the response rate of 67%, which is relatively high compared to previous surveys among pancreatologists [39-41]. Furthermore, the ratio of 30:70 between academic and nonacademic pancreatologists attributed to a valuable insight into the pancreatologists' opinions on the use of therapeutic anticoagulation. This study also has several limitations. First, the results may not directly reflect the actual practice in other countries as only members of two Dutch associations of pancreatology were invited. This decision was made to avoid selection based on publication record, and consequently include pancreatologists who are not actively involved in the treatment of acute pancreatitis[40]. Another advantage of our method is that it allowed us to calculate the survey's response rate by bypassing the confidentiality of membership lists of international pancreatic associations. Second, the clinical presentation of SVT is very heterogeneous, as well as the patient characteristics and clinical disease course among acute pancreatitis patients, which influences current decision making. For this reason, it might have been difficult for pancreatologists to answer some of the general questions. Therefore, case vignettes were used to explore what considerations underpin their decisions. As the descriptions throughout the case vignettes were consistently formulated and only one clinical detail was changed at a time, treatment of patients with superior mesenteric vein and splenic vein thrombosis was not assessed in the case vignettes. Consequently, the pancreatologists' preference on this manifestation of SVT in acute pancreatitis remained unknown. Finally, the rationale behind the "nonprescribing trend" was not assessed adequately, which could be a focus for future research.

#### CONCLUSION

In conclusion, this national survey demonstrates the tendency of pancreatologists to prescribe therapeutic anticoagulation for acute thrombosis, in particular for acute portal vein thrombosis and in case of thrombus progression, irrespective of the presence of infected necrosis. With therapeutic anticoagulation, the majority of pancreatologists believed that the clinical outcomes of acute pancreatitis patients with splanchnic vein thrombosis will improve. Furthermore, this study reflects on several knowledge gaps in literature, and sets out clear points for future research. Specifically, a deeper understanding of the pathophysiology and natural course of splanchnic vein thrombosis secondary to acute pancreatitis would allow us to clarify the therapeutic role of anticoagulation.

#### ARTICLE HIGHLIGHTS

#### Research background

Splanchnic vein thrombosis (SVT) is a severe complication of acute pancreatitis that may cause portal hypertensive complications and bowel ischemia. To prevent such complications, therapeutic anticoagulation is recommended in the general population of patients with an acute SVT.

#### Research motivation

Evidence to support this recommendation in acute pancreatitis patients does however not exist and as a result, clinical decision-making is mostly based on the preferences and beliefs of the pancreatologists.

#### Research objectives

To gain insight into current opinions on the use of therapeutic anticoagulation for SVT in acute pancreatitis.

#### Research methods

An online survey was sent to 139 Dutch pancreatologists. The threshold to assume agreement was set at

#### Research results

The response rate was 67% (n = 93). Seventy-one pancreatologists (77%) regularly prescribed therapeutic anticoagulation for SVT, using LMWH in the acute phase (87%). The majority favored therapeutic anticoagulation for acute thrombosis (90%), portal vein thrombosis in patients with or without infected necrosis (82% and 90%) and in case of thrombus progression (88%). There was no agreement whether the risk of bleeding is a barrier for initiation of therapeutic anticoagulation.

#### Research conclusions

The pancreatologists reached agreement regarding the use of therapeutic anticoagulation for SVT, particularly in cases of acute thrombosis, portal vein thrombosis and thrombus progression.

#### Research perspectives

To get a better understanding of the therapeutic role of anticoagulation, it is crucial to conduct prospective studies targeting the pathophysiology and natural course of SVT in acute pancreatitis patients.

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#### **FOOTNOTES**

Author contributions: Sissingh NJ, Groen JV, and Mieog JSD designed the study; all authors critically assessed the study design; Boekestijn B and Bollen TL provided the radiological images; Sissingh NJ, van Hooft JE, Mieog JSD, and van Eijck CHJ sent or promoted the study; Sissingh NJ did the statistical analysis and wrote the initial draft of the manuscript; Groen JV, Timmerhuis HC, Besselink MG, Boekestijn B, Bollen TL, Bonsing BA, Klok FA, van Santvoort HC, Verdonk RC, van Eijck CHJ, van Hooft JE, and Mieog JSD critically assessed and edited the manuscript; Sissingh NJ coordinated the writing process and revised the manuscript; and all authors read and approved the final manuscript.

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