### Topical hemostatic agents in the management of upper gastrointestinal bleeding: a meta-analysis



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#### **Authors**

Ali A. Alali<sup>1</sup>, Sarvee Moosavi<sup>2</sup>, Myriam Martel<sup>3</sup>, Majid Almadi<sup>4</sup>, Alan N. Barkun<sup>5</sup>

#### Institutions

- 1 Department of Medicine, Faculty of Medicine, Kuwait University, Kuwait
- 2 Division of Gastroenterology, Department of Medicine, University of British Columbia, Vancouver, British Columbia, Canada
- 3 Research Institute of the McGill University Health Center, Montreal, Canada
- 4 Department of Medicine, King Saud University, Riyadh, Kingdom of Saudi Arabia
- 5 Division of Gastroenterology, McGill University Health Center, McGill University, Montreal, Canada

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#### **Corresponding author**

Dr. Alan N. Barkun, McGill University and the McGill University Health Centre, 1650 Cedar Avenue, D7.346, Montréal, QC H3G1A4, Canada alan.barkun@muhc.mcgill.ca

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#### ABSTRACT

Background and study aims Novel topical hemostatic agents have shown promising results in treating patients with non-variceal upper gastrointestinal bleeding (NVU-GIB). However, data are limited even in published metaanalyses as to their role, especially compared to conventional endoscopic approaches. The aim of this study was to perform a highly comprehensive systematic review assessing the effectiveness of topical hemostatic agents in UGIB in different clinical settings.

Methods We performed a literature search of OVID MED-LINE, EMBASE, and ISI Web of Knowledge databases through September 2021. Studies assessing the efficacy of topical hemostatic agents in UGIB were included. Main outcomes were immediate hemostasis and overall rebleeding. Results A total of 980 citations were identified and 59 studies with a total of 3,417 patients were included in the analysis. Immediate hemostasis was achieved in 93% (91%; 94%), with similar results according to etiology (NVUGIB vs. variceal), topical agent used, or treatment strategy (primary vs. rescue). The overall rebleeding rate was 18% (15%; 21%) with the majority of rebleeds occurring in the first 7 days. Among comparative studies, topical agents achieved immediate hemostasis more often than standard endoscopic modalities (OR 3.94 [1.73; 8.96), with non-different overall rebleeding odds (OR 1.06 [0.65; 1.74]). Adverse events occurred in 2% (1%; 3%). Study quality was overall low to very low.

**Conclusions** Topical hemostatic agents are effective and safe in the management of UGIB with favorable outcomes when compared to conventional endoscopic modalities across a variety of bleeding etiologies. This is especially true in novel subgroup analyses that assessed immediate hemostasis and rebleeding among RCTs and in malignant bleeding. Due to methodological limitations of available data, additional studies are needed to ascertain their effectiveness more confidently in the management of patients with UGIB.

#### Introduction

Upper gastrointestinal bleeding (UGIB) is a common emergency condition associated with significant morbidity and mortality [1]. In the United States, UGIB results in more than 300,000 hospital admissions annually with significant associated costs [2, 3]. Despite recent advances in the endoscopic management of both non-variceal upper gastrointestinal bleed (NVUGIB) and variceal upper gastrointestinal bleed, some patients fail conventional endoscopic therapy. Indeed, approximately 15% of patients treated for UGIB fail primary endoscopic therapy, and re-bleeding can occur in up to 25% of patients after initial successful therapy [4] with associated increased morbidity and mortality, especially in patients with variceal bleeding [5].

Recently, novel topical endoscopic hemostatic agents were introduced for the management of UGIB [6]. These agents have shown promising results as a primary or salvage therapy while requiring less technical expertise [6]. Such agents include, among others the hemostatic powder TC-325 (Hemospray, Cook Medical, Bloomington, Indiana, United State), the starch derived polysaccharide hemostatic system (EndoClot, Endoclot Plus Inc., Santa Clara, California, United States), the biocompatible natural polymer UI-EWD (Nexpowder, NextBiomedical Co, Incheon, South Korea) and more recently the synthetic self-assembling peptide agent (PuraStat, 3D-Matrix, Europe Ltd., France) [7,8] (**► Table 1**). Previous studies have concluded that these hemostatic agents are effective and safe in the treatment of NVUGIB with low adverse event (AE) rates [9– 12]. However, reported re-bleeding rates have been high (19% at 72 hours) when used to treat NVUGIB [7]. As a result, the international guidelines for the management of ulcer bleeding suggested the hemostatic powder Tc-325 be used as temporizing measure when conventional endoscopic therapy fails but suggested against using it as a monotherapy [13]. In contradistinction, the more recent American College of Gastroenterology guidelines recently endorsed such a use [14], bolstered by a recently published randomized trial exhibiting potentially limited external validity [15, 16]. Unfortunately, preventing an adequate characterization of overall effectiveness, many of the published studies and systematic reviews did not completely assess the role of Tc-325 according to bleeding etiology which is thought to be a critical consideration when opting for such therapy. [16] Moreover, the published meta-analyses also failed to address other topical hemostatic agents in the management of UGIB or in certain situations such as variceal bleeding, even if data addressing these are sparse.

The aim of this systematic review and meta-analysis is thus to more completely characterize the effectiveness and safety of hemostatic agents in achieving and maintaining hemostasis when managing patients with varying etiologies of UGIB in order to better and more comprehensively guide clinician decisions and future societal recommendations.

Agent	Trade Name	Composition	Mechanism of action	Approved human application	Formulation
TC-325	Hemospray (Cook Medical, Win- ston-Salem, North Car- olina, USA)	Granular mineral- based	Absorbs H <sub>2</sub> O Forms mechanical tamponade Activates clotting cas- cade	Nonvariceal gastroin- testinal bleed	CO <sub>2</sub> pressurized handler canister (20 g)
EndClot	EndoClot (Endoclot Plus Inc., Santa Clara, California, United States)	Absorbable starch- based modified poly- saccharide	Absorbs H <sub>2</sub> O and con- centrates cells Activates clotting cas- cade	Upper and lower gas- trointestinal bleeding	Pressurized air compressor
PuraStat	PuraStat (3D-Matrix, Europe Ltd., France)	Synthetic self-assem- bling peptide agent	Forms a gel coat that induces hemostasis Promotes healing	Bleeding from capillar- ies and oozing from ca- pillaries of the gastro- intestinal tract Prevention of delayed bleeding post colonic ESD	Prefilled syringe
UI-EWD	Nexpowder (NextBiomedical Co, Incheon, South Korea)	Biocompatible natural polymer	Forms adhesive hydro- gel in the presence of water	Intended for upper gastrointestinal bleed- ing	Spraying device
CEGP-003	CEGP-003 (CGBio, Seongnam, South Korea)	Absorbable and adhe- sive macromolecules containing epidermal growth factor (EGF)	Forms adhesive gel upon contact with moist mucosa facilitat- ing hemostasis Promotes wound heal- ing	Treatment of upper gastrointestinal bleed- ing Prevention of bleeding post EMR/ESD	Spraying device

**Table 1** Summary of included topical hemostatic agents.

ESD, endoscopic submucosal dissection; EMR, endoscopic mucosal resection.

### Methods

#### Search strategy

A comprehensive literature search was performed, from the recorded start of databases to September 2021 using OVID MED-LINE, EMBASE, and ISI Web of Knowledge databases, with a combination of MeSH term and controlled vocabulary to identify studies related to: 1) hemostatic agent or powder and; 2) gastrointestinal bleeding (Supplementary Table 1). Abstracts presented at major gastroenterology conferences (ACG, CDDW, DDW, UEGW, APDW) in the past 5 years were also hand-searched. Additional relevant studies were identified from cross-referencing and hand-searches of references of retrieved articles.

# Validity assessment, data abstraction and rating of evidence

Two reviewers (AA, MM) evaluated the eligibility of all identified citations independently, with a third (AB) resolving disagreements. Study quality was assessed using the Cochrane Risk of bias tool for randomized trials [17], and the Ottawa-Newcastle criteria for observational studies [18]. We used the GRADE rating to characterize the certainty of evidence [19].

#### Inclusion and exclusion criteria

Studies assessing any hemostatic agent in UGIB of any etiology were considered for inclusion. Both manuscripts and abstracts from major gastroenterology societies (limited to the previous 5 years) were considered for inclusion. Only studies published in English were included. We excluded case reports, studies with less than 10 patients, review articles, and non-human studies. In addition, we excluded studies reporting on the hemostatic agent "Ankaferd" due to its limited availability globally [6]. Studies that used hemostatic agents prophylactically to prevent GI bleeding were also excluded.

#### Study definitions

"Immediate hemostasis" was defined as no further bleeding at least 3 minutes after application of the hemostatic agent. "Rebleeding" was defined as evidence of recurrent UGIB manifested as overt gastrointestinal bleeding or drop in hemoglobin≥2g/dL after achieving immediate hemostasis [20]. Rebleeding data were included only for patients following initial immediate hemostasis. "Technical success" was defined as successful use of the hemostatic agent during endoscopy without any technical problems (e.g. blockage of applications catheter). "Monotherapy" was defined as the use of a hemostatic agent alone, whereas "combination therapy" was defined as using the hemostatic agent in combination with adjunctive conventional endoscopic methods. "Rescue therapy" was defined as the use of a hemostatic agent when other conventional endoscopic methods had failed as evidenced by failure of immediate hemostasis. "Primary therapy" was defined as using the hemostatic agent as first-line endoscopic therapy for bleeding.

#### Outcomes

Immediate hemostasis and overall rebleeding (defined as any rebleeding during follow-up after the index endoscopy) were the outcomes of interest. Other outcomes assessed included 7-day and 30-day rebleeding, overall-mortality, bleeding-related mortality, technical success, and AEs related to application of the topical hemostatic agent. Data will be present in turn for a meta-analysis of proportions (purely descriptive) and in a meta-analysis assessing studies that compared hemostatic powders to a control therapeutic approach.

#### Sensitivity and subgroup analyses

Pre-planned possible subgroup and sensitivity analyses included assessments according to: type of hemostatic agent, type of therapies (primary therapy, rescue therapy, monotherapy, combination therapy), type of lesions (peptic ulcer disease (PUD), post endoscopic intervention, varices, malignant lesions), randomized-controlled trials alone; fully published articles alone, higher quality studies, year of publication, continent where the study was performed, performing a fixed rather than a random effect model (when appropriate), and when correcting for double-zero events.

#### Statistical analysis

Categorical estimates of outcomes were reported as proportions and 95% confidence intervals (CI) using weighted random effects models. Continuous variables are reported as means and standard deviations medians were used if means were not available and standard deviations (SDs) were calculated or imputed when possible. For comparative studies, effect size was calculated with weighted mean differences (WMDs) for continuous variables. Odds ratios (ORs) are expressed for categorical variables. The DerSimonian and Laird method for random effect models was applied to all outcomes to determine corresponding overall effect sizes and their confidence intervals. Sensitivity analyses were performed using the Mantel-Haenszel method with random effect models; however fixed effects models were used when no statistical heterogeneity was noted. WMD were handled as continuous variables using the inverse variance approach. Presence of heterogeneity across studies was defined using a Chi-square test of homogeneity with a 0.10 significance level. The Higgins I<sup>2</sup> statistic was calculated to quantify the proportion of variation in treatment effects attributable to between-study heterogeneity, with values of 25%, 50%, and 75% representing low, moderate, and high heterogeneity, respectively. When heterogeneity was noted, prediction intervals were calculated and added to the forest plot. The prediction interval calculates the 95% of where the effect size will be if a new study is randomly added to the meta-analysis. In order to ensure that zero event trials did not significantly affect the heterogeneity or p-values, sensitivity analyses were performed where a continuity correction was added to each trial with zero events using the reciprocal of the opposite treatment arm size. For all comparisons. Publication bias was evaluated using funnel plots if at least three citations were identified. All statistical analyses were done using Revman 5.4 and Meta package in R version 2.13.0, (R Foundation for Statistical Computing, Vienna, Austria, 2008).

#### Results

# Included studies, quality assessment and publication bias

We initially identified 980 citations. After review of abstracts, a total of 59 studies were included (**> Fig. 1**). One study was excluded since it reported results as per episode of bleeding rather than per patient outcomes [21].

For the outcomes in the (solely descriptive) meta-analysis of proportions, moderate to high heterogeneity was present for immediate hemostasis, rebleeding (overall, 7 days and 30 days) and overall mortality. No significant heterogeneity was noted in the portion of the meta-analysis addressing betweengroup comparative results. No publication bias was observed (data available upon request).

Study quality was overall low-very low with the Ottawa Newcastle Scores (NOS) ranging from 4 to 8 stars out of a possible score of 9, with a mean of  $5.9\pm0.8$  for observational studies. Assessing the individual domains of the NOS confirmed the low studies' quality (Supplementary Table 2). The Cochrane risk bias tool revealed a high potential for performance bias across studies for randomized trials since all were single-blinded (Supplementary Fig. 1). The grading of the evidence was performed in studies that included a comparative arm and was found to be "very low" for all outcomes (Supplementary Table 3).

#### Patient and study characteristics

Overall, 59 studies were included (n=3417 patients). Hemospray (TC-325) was the sole intervention in 44 studies [15, 22–64], EndoClot in five [65–69], PuraStat in four [70–73], Nexpowder in three [74–76] and CEGP-003 [77] in one. In addition, two studies included both Hemospray and EndoClot [78, 79]. Seven were RCTs, and 15 were prospective while 37 were retrospective cohort studies. Furthermore, 44 were fully published articles, while the remainder were sole abstracts (**► Table 2**). Ten studies included a comparison with standard endoscopic therapy [15, 24, 30, 39, 42, 47, 52, 65, 69, 77]. Reports recruited patients between 2009 and 2021. Most of the bleeding lesions were classified as oozing bleeding (Forrest Ib). There were 11 studies [26–28, 37, 38, 49, 50, 57, 60, 64, 78] (n=124 patients) that addressed patients with variceal bleeding. Full study and patient characteristics are shown in **► Table 2**.

#### Immediate hemostasis and overall rebleeding

Immediate hemostasis was reported in 59 studies (n = 2,919 patients) with a pooled success rate of 93% (91%; 94%) ( $\triangleright$  Fig. 2,  $\triangleright$  Table 3). Overall rebleeding was included in 58 studies (n = 2,696) with a pooled overall rebleeding rate of 18% (15%; 21%) ( $\triangleright$  Fig. 3,  $\triangleright$  Table 3). Rebleeding rates at 7 and 30 days were 17% (14%; 20%) and 21% (17%; 27%), respectively ( $\triangleright$  Table 3).



▶ Fig. 1 PRISMA diagram of included studies. From: *Page MJ*, *McKenzie JE*, *Bossuyt PM* et al. The PRISMA 2020 statement: an updated guideline for reporting systematic reviews. BMJ 2021; 372: n71.

In the 10 studies comparing the topical hemostatic agent to a control intervention (n=797), immediate hemostasis was more significantly achieved with the former (odds ratio [OR] = 3.94 (1.73; 8.96) (**Table 4** and **Fig. 4**). Among the comparative studies, the overall rebleeding risk was not significantly different between topical agents and conventional endoscopic therapy (OR = 1.06; 0.65–1.74, 10 studies, n=775); this was also the case for rebleeding at 7 and 30 days (OR = 0.97; 0.43– 2.16) and OR = 0.75; 0.39–1.45, respectively) (**Table 4** and **Fig. 5**).

#### Other outcomes

Pooled proportions for overall mortality (45 studies, n = 2,245) and bleeding-related mortality (34 studies; n = 1,563) were 15% (12%; 19%) and 5% (4%; 7%), respectively. In terms of technical success (52 studies, n = 2,392) and AEs (45 studies, n = 2,111), analyses yielded pooled proportions of 97% (96%; 98%) and 2% (1%; 3%), respectively.

In studies that included a comparison group, none of these outcomes differed significantly when comparing the topical agent to the control hemostatic modality(ies) (table 4).

Study Country	Interven- tion	Compari- son	Sample size	Design	Study period	Indication	Etiology	Bleeding severity
Arena et al. 2017 [23] Italy	Hemospray	None	15	Retrospective cohort	2014– 2015	Primary therapy	Malignancy (100%)	Oozing (100%)
Appleby et al. 2017 [22] UK Abstract	Hemospray	None	32	Retrospective cohort2014- 2017Primary therapyPUD (56%) Malignancy (9%) Rescue therapy (28%) Others (7%)		NR		
Baracat et al. 2020 [24] Brazil	Hemospray	Hemoclips	39	RCT 2015- Primary PUD (43.6 2017 therapy Malignance Post interv (10.3%) Dieulafoy I (5.1%) MalloryWe (7.7%) Others (20		PUD (43.6%) Malignancy (12.8%) Post intervention (10.3%) Dieulafoy lesion (5.1%) MalloryWeiss tear (7.7%) Others (20.5%)	Spurting (10.3 %) Oozing (89.7 %)	
Bestari et al. 2020 [26] Indonesia	Hemospray	None	30	Retrospective 2016– cohort 2019		Primary therapy Rescue therapy	PUD (56.7%) Malignancy (26.7%) Variceal bleed (10%) Portal hypertensive gastropathy (6.7%)	NR
Becq et al. 2020 [25] France	Hemospray	None	152	Retrospective cohort	trospective 2015– Primary Pl hort 2018 therapy M Rescue Es therapy O		PUD (47.7%) Malignancy (22.2%) Esophagitis (12.4%) Other (17.7%)	Oozing (84.3%) spurting (11.1%) None active (2.6%) Undetermined (2.0%)
Chahal et al. 2020 [28] Canada	Hemospray	None	73	Retrospective cohort	2014- 2018	Primary therapy Rescue therapy	PUD (67.1%) Malignancy (5.5%) Variceal bleed (5.5%) MWT (8.2%) Other (13.7%)	For PUD (n = 49) Forrest la (14.3 %) Forrest lb (53.1 %) Forrest lla (20.4 %) Forrest llb (14.3 %) Forrest lll (4.1 %)
Cahyadi et al. 2017 [27] Germany	Hemospray	None	52	Retrospective cohort	2013– 2017	Primary therapy Rescue therapy	PUD (34.6%) Malignancy (19.2%) Post procedure (30.7%) Dieulafoy lesion (1.9%) GAVE (1.9%) Variceal bleed (3.8%) Other (7.6%)	Peptic ulcer only, n = 18 Forrest Ib (38.9%) Forrest IIa (33.3%) Forrest IIb (22.2%) Forrest IIc (5.6%)
Chen et al. 2015 [29] Canada	Hemospray	None	56	Retrospective cohort	2011- 2013	Primary therapy Rescue therapy	Benign NVUGIB (37.5%) Malignancy (33.9%) Post intervention (28.6%)	For PUD (n = 13) Forrest Ia (23.1%) Forrest Ib (76.9%)
Chen et al. 2020 [30] Canada	Hemospray	СНР	17	RCT	2014– 2016	Primary therapy	Malignancy (100%)	Spurting (5%) Oozing (95%)

#### **Table 2** Patient and study characteristics.

► Table 2 (Continuation)										
Study Country	Interven- tion	Compari- son	Sample size	Design	Study period	Indication	Etiology	Bleeding severity		
De Santiago et al. 2019 [60] Spain	Hemospray	None	219	Retrospective cohort	2011– 2018	Primary therapy Rescue therapy	PUD (33.3%) Malignancy (21.0%) Post-procedure (14.2%) postsurgical (6.4%) Dieulafoy lesion (3.7%) Variceal bleed (3.2%) GAVE (0.9%) Others (17.4%)	Peptic ulcer only, n = 73 Forrest Ia (24.7 %) Forrest Ib (64.4 %) Forrest IIb (65.5 %) Forrest IIb (5.5 %)		
Disney et al. 2015 [31] UK Abstract	Hemospray	None	19	Retrospective cohort	NR	Primary therapy Rescue therapy	Malignancy (100%)	Oozing (100%)		
Giles et al. 2016 [32] New Zeland	Hemospray	None	36	Retrospective cohort	2013- 2016	Rescue Therapy	PUD (66.7%) Malignant (5.6%) post intervention (16.7%) Others (11.0%)	Spurting (13.9%) Oozing (69.4%) Visible vessel (5.6%) Adherent clot (5.6%) Clean base (5.6%)		
Gupta et al. 2018 [33] UK Abstract	Hemospray	None	45	Retrospective cohort	2013- 2017	Primary therapy Rescue therapy	PUD (41%) Malignancy (15%) Post-intervention (15%) Other (29%)	Forrest Ia (15%) Forrest Ib (46%) Forrest IIa (15%) Forrest IIb (15%) Forrest IIc (9%)		
Haddara et al. 2016 [34] France	Hemospray	None	202	Retrospective cohort	2013- 2015	Primary therapy Rescue therapy	PUD (37.1%) Malignant (30.2%) Post-intervention (17.3%) Dieulafoy lesion (1.5%) Others (3.9%)	For PUD (n = 75) Forrest Ia (20.0%) Forrest Ib (57.3%) Forrest II (18.7) Unclassified (4.0%)		
Hagel et al. 2017 [35] Germany	Hemospray	None	33	Retrospective cohort	2013- 2014	Rescue Therapy	PUD (48.5%) Malignancy (12.1%) Post intervention (12.1%) Diffuse bleeding (18.2%) Others (9.1%)	NR		
Holster et al. 2015 [36] Netherland	Hemospray	None	16	Retrospective cohort	2011- 2012	Primary therapy Rescue therapy	PUD (56.0%) Malignant (13.0%) Others (31.0%)	For PUD (n = 9) Forrest Ia (55.5%) Forrest Ib (44.4%)		
Hussein et al. 2021 (1) [62] UK	Hemospray	None	105	Prospective study	2016- 2020	Primary therapy Rescue therapy	Malignancy (100%)	Spurting (6%) Oozing (77%) Visible vessel/adher- ent clot (12%)		
Hussein et al. 2021 (2) [61] UK	Hemospray	None	202	Prospective cohort	2016- 2019	Primary therapy Rescue therapy	PUD (100%)	Forrest Ia (19.0%) Forrest Ib (58.0%) Forrest IIa (12.0%) Forrest IIb (10.0%)		
Hussein et al. 2021 (3) [64] UK Abstract	Hemospray	None	12	Prospective study	2016– 2019	Primary therapy Rescue therapy	Esophageal varices (83.3%) Gastric varices (16.7%)	NR		

Study Country	Interven- tion	Compari- son	Sample size	Design	Study period	Indication	Etiology	Bleeding severity
Hussein et al. 2020 [63] UK	Hemospray	None	73	Prospective cohort	2016- 2019	Primary therapy Rescue therapy	Post-procedure (100%)	NR
Ibrahim et al. 2015 [37] Belgium	Hemospray	None	30	Prospective cohort2013- 2014Primary TherapyEsophageal varices (83.4%) Gastric varices (10%) Duodenal varices (6.6%)		Spurting (43.4%) Fibrin plug/red streaks (56.6%)		
Ibrahim et al. 2019 [38] Belgium	Hemospray	Early elec- tive endos- copy	86	RCT2014- 2016Primary therapyVariceal bleed (100%)		Variceal bleed (100%)	Spurting (16.3%) Blood in stomach (83.7%)	
Kwek et al. 2017 [39] Singapore	Hemospray	Convention- al endo- scopic ther- apy	20	RCT	2013– Primary PUD (100%) 2015 therapy		PUD (100%)	Forrest Ia (5.0%) Forrest Ib (35.0%) Forrest IIa (50.0) Forrest IIb (10.0%)
Lau et al. 2020 [15] China	Hemospray	Convention- al endo- scopic ther- apy	224	RCT 2015- Primary PUD (58%) 2018 therapy Malignancy (1 Others (27.75		PUD (58 %) Malignancy (14.3 %) Others (27.7 %)	NR	
Leblanc et al. 2013 [40] France	Hemospray	None	17	Retrospective cohort2011- 2012Primary therapy Rescue therapyPost intervention (70.6%) Malignancy (29.4%)		For post-intervention (n = 12) pulsatile (8.3 %) oozing (91.7 %)		
Malik et al. 2015 [41] UK Abstract	Hemospray	None	19	Retrospective cohort	2013- 2014	Primary therapy Rescue therapy	PUD (53%) Malignancy (21%) Post intervention (21%) Other (5%)	For PUD (n = 10) Forrest la (0%) Forrest lb (60%) Forrest lla (30%) Forrest llb (10%)
Martins et al. 2019 [42] Brazil Abstract	Hemospray	СНР	36	RCT	2016– 2017	Primary therapy	Malignancy (100%)	Active bleeding (63.9%)
Masci et al. 2014 [43] Italy	Hemospray	None	13	Prospective cohort	NR	Primary therapy	PUD (100%)	Forrest Ia (35.7 %) Forrest Ib (64.3 %)
Meng et al. 2018 [44] Canada	Hemospray	None	25	Retrospective study	2010– 2016	Primary therapy Rescue therapy	Malignancy (100%)	Forrest Ia (8.0%) Forrest Ib (76.0%) unclassified (16.0%)
Min et al. 2018 [45] UK Abstract	Hemospray	None	48	Retrospective cohort	ective 2016– Primary PUD (69.0%) 2017 therapy Malignancy (2. Rescue Post- intervent therapy (4.0%) GAVE (4.0%) Esophagitis (4. Others (16.0%)		PUD (69.0%) Malignancy (2.0%) Post- intervention (4.0%) GAVE (4.0%) Esophagitis (4.0%) Others (16.0%)	NR
Nasr et al. 2015 [46] UK Abstract	Hemospray	None	26	Retrospective cohort	2013– 2015	Primary therapy Rescue therapy	NVUGIB (65.4%) Post-intervention (34.6%)	NR

► Table 2 (C	ontinuation)							
Study Country	Interven- tion	Compari- son	Sample size	Design	Study period	Indication	Etiology	Bleeding severity
Paoluzi et al. 2021 [79] Italy	Hemospray EndoClot	СНР	108	Retrospective cohort	2017- 2019	Primary therapy Rescue therapy	PUD (66.7%) Malignancy (21.3%) Post-intervention (6.4%) Others (5.5%)	For PUD (n = 72) Forrest IA (20.8%) Forrest IB (79.2%) For malignancy (n = 23) oozing (100%)
Pittayanon et al. 2016 [47] Thailand	Hemospray	СНР	20	Prospective cohort	2014– 2015	Primary therapy	Malignancy (100%)	oozing blood (100%
Pittayanon et al. 2018 [48] Canada	Hemospray	None	79	Retrospective cohort	2011– 2016	Primary therapy Rescue therapy	Malignancy (100%)	Adherent clot (4.5%) Blood oozing (94.3%) Blood spurting (1.1%)
Prentice et al. 2018 [49] UK Abstract	Hemospray	None	47	Retrospective cohort	2014– 2017	Primary therapy Rescue therapy	NVUGIB (78.7%) Variceal bleed (21.3%)	NR
Ramirez-Polo et al. 2019 [50] Mexico	Hemospray	None	81	Retrospective cohort	2015– 2017	Primary therapy	PUD (17.3%) Malignancy (43.2%) post-procedure (14.8%) Dieulafoy lesion (6.1%) Variceal bleed (2.5%) Postsurgical (2.5%) Other (13.6%)	NR
Shivaji et al. 2018 [51] UK Abstract	Hemospray	None	45	Retrospective cohort	2013- 2017	Primary therapy Rescue therapy	PUD (41.0%) Malignancy (15.0%) Post-intervention (15.0%) Others (29.0%)	Peptic ulcer only, n = 22 Forrest la (15.0%) Forrest lb (46.0%) Forrest lla (15.0%) Forrest llb (15.0%) Forrest llc (9.0%)
Sinha et al. 2016 [52] UK	Hemospray	СНР	40	Retrospective cohort	2013- 2015	Rescue therapy	PUD (65.0%) Esophageal ulcer (20%) Others (15.0%)	Forrest Ia (60.0%) Forrest Ib (40.0%)
Smith et al. 2014 [53] UK	Hemospray	None	63	Prospective cohort	2011	Primary therapy Rescue therapy	PUD (57.1%) Malignancy (12.6%) Post-procedure (15.9%) Dieulafoy lesion (3.2%) MWT (3.2%) GAVE (3.2%) Other (5.8%)	Peptic ulcer only, n = 30 Forrest la (36.7 %) Forrest lb (53.3 %) Unclassified (10 %)
Sulz et al. 2014 [54] Switzerland	Hemospray	None	15	Prospective cohort	2013	Primary therapy Rescue therapy	PUD (25.0%) Malignancy (18.8%) Post-intervention (12.5%) Others (43.8%)	Spurting (18.8%) oozing (81.2%)
Sung et al. 2011 [55] Hong Kong	Hemospray	None	20	Prospective cohort	2009– 2010	Primary Therapy	PUD (100%)	Forrest Ia (5%) Forrest Ib (95%)

Study Country	Interven- tion	Compari- son	Sample size	Design	Study period	Indication	Etiology	Bleeding severity
Thayalasekaran et al. 2017 [56] UK Abstract	Hemospray	None	44	Retrospective cohort	Retrospective 2014– Primary PUD (65.9%) cohort 2016 therapy Malignancy (2.3%) Rescue Variceal bleed therapy (6.8%) MWT (4.5%) Post-intervention (12.3%) Others (8.2%)		PUD (65.9%) Malignancy (2.3%) Variceal bleed (6.8%) MWT (4.5%) Post-intervention (12.3%) Others (8.2%)	Forrest Ia (26.5%) Forrest Ib (44.1%) Forrest IIa (23.5%) Forrest IIb (5.9%)
Vitali et al. 2019 [78] Germany	Hemospray	EndoClot	127	Prospective cohort	ospective 2013– Primary PUD (38.6%) hort 2017 Primary Malignancy ( Rescue Variceal bleed therapy (10.2%) Reflux esoph (9.4%) Others (30.0		PUD (38.6%) Malignancy (11.8%) Variceal bleed (10.2%) Reflux esophagitis (9.4%) Others (30.0%)	NR
Weaver et al. 2019 [57] USA Abstract	Hemospray	None	12	Prospective 2018– Pri cohort 2019 the Rei the		Primary therapy Rescue therapy	PUD (50.0%) MWT (16.7%) Variceal bleed (8.3%) GAVE (8.3%) Post-intervention (16.7%)	Peptic ulcer only, n = 6 Forrest Ia (50.0%) Forrest Ib (50.0%)
Widlak et al. 2015 [58] UK Abstract	Hemospray	None	48	Retrospective cohort	pective 2013– Primary 2015 therapy Rescue therapy		PUD (60.5%) Other (27%) Post banding ulcer (12.5%)	NR
Yau et al. 2014 [59] Canada	Hemospray	None	19	Retrospective cohort	2012- 2013	Primary therapy Rescue therapy	PUD (63.2%) Dieulafoy lesion (10.5%) Mucosal erosion (5.3%) Angiodysplasia (5.3%) Post-intervention (10.6%) Unknown (5.3%)	Spurting (21.1%) oozing (57.9%) No active bleeding (21.1%)
Branchi et al. 2021 [72] Germany	PuraStat	None	78	Prospective cohort	2017– 2018	Primary therapy Rescue therapy	PUD Post-intervention Malignancy Vascular lesions	spurting (6%) Oozing (69%) Visible vessel (14%) Adherent clot (5%) Flat pigmented spot (5%)
de Nucci et al. 2020 [70] Italy	PuraStat	None	41	Retrospective cohort	2017– 2019	Rescue therapy	PUD (46.3%) Malignancy (7.3%) Post-intervention (39.0%) Other (7.4%)	Spurting (19.5%) Oozing (80.5%)
Labianca et al. 2021 [73] Italy Abstract	PuraStat	None	15	Retrospective cohort	2018– 2020	Primary therapy Rescue therapy	PUD (86.7%) Post-intervention (13.3%)	NR
Subramaniam et al. 2019 [71] UK	PuraStat	None	44	Retrospective cohort	2016– 2017	Primary therapy	Post-intervention (100%)	NR

► Table 2 (Continuation)										
Study Country	Interven- tion	Compari- son	Sample size	Design	Study period	Indication	Etiology	Bleeding severity		
Beg et al. 2015 [65] UK	EndoClot	СНР	130	Retrospective 2012– cohort 2014		Rescue therapy	PUD (90%) Esophageal ulcer (4.6%) MWT (3.8%) Other (1.6%)	NR		
Hagel et al. 2020 [68] Germany	EndoClot	None	22	Retrospective cohort	2015- 2020	Primary therapy Rescue therapy	PUD (40.9%) Malignancy (40.9%) Others (18.2%)	NR		
Kim et al. 2018 [66] South Korea	EndoClot	None	12	Retrospective cohort	2016– 2017	Primary therapy	Malignancy (100%)	Forrest lb (100%)		
Park et al. 2019 [69] South Korea	EndoClot	СНР	176	Retrospective cohort	2012- 2017	Primary therapy Rescue therapy	PUD (68.8%) Malignancy (21.6%) Post-intervention (5.7%) Radiation gastritis (3.9%)	Forrest Ia (9.7 %) Forrest Ib (54.5 %) Forrest IIa (35.8 %)		
Prei et al. 2016 [67] Germany	EndClot	None	58	Prospective cohort	2012- 2014	Primary therapy Rescue therapy	PUD (46.6%) Malignancy (17.2%) Post-intervention (10.3%) Esophagitis (10.3%) Other (15.6%)	Forrest Ia (5.2%) Forrest Ib (65.5%) Forrest IIa (10.3%) Forrest IIb (6.9%) Forrest IIc (12.1%)		
Park et al. 2019 (1) [74] South Korea	Nexpowder	None	17	Prospective cohort	2016- 2017	Rescue therapy	PUD (29.4%) Malignancy (23.5%) Post-intervention (41.2%) Other (5.9%)	Forrest Ia (11.8%) Forrest Ib (88.2%)		
Park et al 2019 (2) [75] South Korea Abstract	Nexpowder	None	56	Retrospective cohort	NR	Primary therapy	PUD (14.3%) Malignancy (1.8%) Post-intervention (82.1%) Other (1.8%)	NR		
Shin et al. 2021 [76] South Korea	Nexpowder	None	41	Retrospective cohort	2016– 2019	Primary therapy Rescue therapy	Malignancy (100%)	Forrest Ia (7.3 %) Forrest Ib (92.7 %)		
Bang et al. 2018 [77] South Korea	CEGP-003	Epinephrine injection	72	RCT	2014– 2015	Primary therapy	PUD (22.2%) Post-intervention (77.8%)	Forrest Ib (83.3%) Forrest IIa (6.9%) Forrest IIb (9.8%)		

PUD, peptic ulcer disease; RCT, randomized controlled trial; CHP, conventional hemostatic procedures; MWT, Mallory-Weiss tearing; GAVE, gastric antral vascular ectasia; NVUGIB, non-variceal upper gastrointestinal bleeding.

#### Sensitivity and subgroup analyses

Results for sensitivity analyses for immediate hemostasis and rebleeding are described in Appendix 1–4. Results confirm the robustness of the findings described in the primary analysis.

When analyzed according to etiology of UGIB, immediate hemostasis rates were high for all etiologies of UGIB including PUD (91% [88%; 93%]) malignancy (94% [91%; 95%]), postendoscopic intervention (90% [85%; 93%]), and a variceal etiology of bleeding (87% [79%; 92%]). The risk of rebleeding for NVUGIB was 22% (18%; 27%), for variceal bleeding was 23% (10%; 45%), and for malignant bleeding, in particular, was 24% (19%; 30%) (Appendix 2).

When grouped according to indication, immediate hemostasis rates were similar when these agents were used as primary or rescue therapy (93%, [89%; 95%] and 90%, [85%; 93%], respectively); these rates varied between 86% to 99% according to the topical agent assessed. The risk of rebleeding was 25% (20%; 30%) when the topical agent was used as rescue therapy, and 18% (14%; 22%) when used as primary therapy.

Study Intervent	tion Tota	al Proportion	95%-Cl	Study	Interventio	on Total	Proportion 95%-Cl
Setting: Hemospray				Setting: CEG	P-003		
Appleby, 2017 28	8 32	0.88	[0.71; 0.96]	Bang, 2018	35	35	1.00 [0.90; 1.00]
Arena, 2016 15	5 15		0.78; 1.00	5,			
Baracat, 2019 19	9 19	1.00	0.82; 1.00]	Setting: Endo	Clot		
Becq, 2020 121	1 152	0.80	0.72; 0.86	Beg, 2015	21	21	1.00 [0.84; 1.00]
Bestari, 2020 30	0 30		0.88; 1.00]	Hagel, 2020	18	22	0.82 [0.60; 0.95]
Cahyadi, 2017 51	1 52	+ 0.98	0.90; 1.00]	Kim, 2018	12	12	
Chahal, 2020 65	5 73		[0.80; 0.95]	Paoluzi, 2021	9	10	0.90 [0.55; 1.00]
Chen, 2015 55	5 56	0.98	[0.90; 1.00]	Park, 2018	36	37	0.97 [0.86; 1.00]
Chen, 2020 7	7 8		[0.47; 1.00]	Prei, 2016	41	58	0.71 [0.57; 0.82]
de Santiago, 2019 203	3 219		[0.90; 0.97]	Vitali, 2019	21	25	0.84 [0.64; 0.95]
Disney, 2015 18	8 19		[0.74; 1.00]	Subtotal		185	0.80 [0.73; 0.86]
Giles, 2016 36	5 36	1.00	[0.90; 1.00]	Random effe	cts model		0.87 [0.75; 0.94]
Gupta, 2018 45	5 45	<u>+</u> 1.00	[0.92; 1.00]	Heterogeneity	y: I <sup>2</sup> = 54%, P	9 = 0.04	
Haddara, 2016 195	5 202		[0.93; 0.99]				
Hagel, 2017 32	2 33	0.97	[0.84; 1.00]	Setting: Pura	stat		
Holster, 2012 13	3 16	0.81	[0.54; 0.96]	Branchi, 2021	68	78	0.87 [0.78; 0.94]
Hussein, 2020 73	3 73	-1.00	[0.95; 1.00]	de Nucci, 202	0 35	41	0.85 [0.71; 0.94]
Hussein, 2021 (1) 102	2 105		[0.92; 0.99]	Labianca, 202	1 15	15	
Hussein, 2021 (2) 178	8 202	0.88	[0.83; 0.92]	Subramaniam	,2019 36	44	0.82 [0.67; 0.92]
Hussein, 2021 (3)	9 12-	• 0.75	[0.43; 0.95]	Subtotal		178	
Ibrahim, 2015 30	0 30	1.00	[0.88; 1.00]	Random effe	cts model		
Ibrahim, 2019 38	8 43	0.88	[0.75; 0.96]	Heterogeneity	y: I <sup>2</sup> = 0 %, P =	= 0.57	
Kwek, 2017 9	9 10		[0.55; 1.00]				
Lau, 2021 108	8 111	0.97	[0.92; 0.99]	Setting: Nex	powder		
Leblanc, 2013 17	7 17		[0.80; 1.00]	Park, 2019 (1)	) 16	17	0.94 [0.71; 1.00]
Malik, 2015 17	7 19		[0.67; 0.99]	Park, 2019 (2)	) 54	56	0.96 [0.88; 1.00]
Martins, 2019 28	8 28	1.00	[0.88; 1.00]	Shin, 2021	40	41	0.98 [0.87; 1.00]
Masci, 2014 13	3 13		[0.75; 1.00]	Subtotal		114	🔶 0.96 [0.91; 0.99]
Meng, 2019 22	2 25	0.88	[0.69; 0.97]	Random effe	cts model		🔶 0.96 [0.91; 0.99]
Min, 2018 46	5 48	0.96	[0.86; 0.99]	Heterogeneity	y: I²= 0 %, P =	= 0.82	
Nasr, 2015 21	1 26	0.81	[0.61; 0.93]				
Paoluzi, 2021 32	2 33	0.97	[0.84; 1.00]	Total		2919	<b>•</b> 0.88 [0.87; 0.89]
Pittayanon, 2016 10	0 10		[0.69; 1.00]	Random effe	cts model		• 0.93 [0.90; 0.94]
Pittayanon, 2018 77	7 79	+	[0.91; 1.00]	Prediction in	terval		[0.72; 0.98]
Prentice, 2018 47	7 47		[0.92; 1.00]	Heterogeneity	y: l²= 67%, P	°<0.01	
Ramirez-polo, 2019 80	0 81		[0.93; 1.00]			0.5	0.6 0.7 0.8 0.9 1
Shivaji, 2018 45	5 45	1.00	[0.92; 1.00]				
Sinha, 2016 19	9 20		[0.75; 1.00]		1.07	10	
Smith, 2014 55	5 63		[0.77; 0.94]	lest for subgr	oup differen	ices (fixe	d effect): $\chi_4^2$ = 18.49, df = 4 ( <i>P</i> < 0.01)
Sulz, 2014 14	4 15		[0.68; 1.00]	lest for subgr	oup differen	ices (ran	dom effects):
Sung, 2011 19	9 20		[0.75; 1.00]	$\chi_4^2 = 15.86$ , df	= 4 (P < 0.01)	)	
Thayalasekaran, 20174	1 44	0.93	[0.81; 0.99]				
Vitali, 2019 68	8 102	0.67	[0.57; 0.76]				
Weaver, 2019 IC		0.83	[0.52; 0.98]				
Viulak, 2015 40	0 48 0 10	0.96	[0.80; 0.99]				
Idu, 2014 Ič	<b>7407</b>		0.74, 1.00]				
Pandom offects model	2407	0.03	0.07, 0.90]				
Heterogeneity: $l^2 = 60\%$	P<0.01	0.93	0.91, 0.95]				
neterogeneity. 1 - 09%	, 1 \0.01						
		0.5 0.6 0.7 0.8 0.9 1					
		.,					

**Fig.2** Forrest plot of primary outcome by proportion: immediate hemostasis.

When studied according to topical agent studied, immediate hemostasis among the different agents was (Hemospray 93% [91%; 95%]), EndoClot (87% [75%; 94%]), PuraStat (86% [80%; 90%]) Nexpowder (96% [91%; 99%]) and CEGP-003 (99% [81%; 100%]).The risk of rebleeding was Hemospray 20% (17%; 24%), EndoClot (10% [6%; 16%]), PuraStat (7% [1%; 27%]) Nexpowder (8% [3%; 20%]) and CEGP-003 (9% [3%; 23%]).

The subgroup analysis of sole RCTs yielded similar outcomes with a proportion of immediate hemostasis of 94% (89%; 97%), and a rebleeding rate of 26% (11%; 48%). Among Sensitivity a-

nalysis using comparative RCTs data confirmed that immediate hemostasis was more frequently achieved with topical hemostatic agents compared to conventional endoscopic therapy (3.62 [1.29; 10.11]), with similar odds of rebleeding (1.27 [0.62;2.59]). Topical agents were more effective when used as primary therapy (3.83 [1.59,9.24]) and when used for malignant lesions (14.74 [2.16,100.61]) (Appendix 3 and 4).

#### **Table 3** Primary and secondary outcome for proportions.

	No. studies	No.patients	Proportion (95 % Cl)	<i>P</i> value for heterogeneity	<b>I</b> <sup>2</sup>
Primary outcome					
Immediate hemostasis (overall UGIB)	59	2919	0.93 (0.91; 0.94)	< 0.01	67%
Overall rebleeding	58	2696	0.18 (0.15; 0.21)	<0.01	69%
Rebleeding 7 days	42	1943	0.17 (0.14; 0.20)	<0.01	55%
Rebleeding 30 days	34	1692	0.21 (0.17; 0.26)	<0.01	75%
Secondary outcome					
Overall mortality	45	2245	0.15 (0.12; 0.19)	<0.01	64%
Bleeding-related mortality	34	1563	0.05 (0.04; 0.07)	0.42	3 %
Technical success	52	2392	0.97 (0.96; 0.98)	0.99	0 %
Adverse events	45	2111	0.02 (0.01; 0.03)	0.99	0 %
LICID upper costraintestinal blooding					

#### Discussion

To our knowledge, this is the largest, most granular and up-todate meta-analysis that has characterized the efficacy and safety of different topical hemostatic agents according to various UGIB etiologies. Results confirm the effectiveness and safety of topical hemostatic agents when used to treat UGIB, however with a low to very low certainty of evidence for the data that these observations are based on, as is also the case for argon plasma coagulation and soft monopolar electrocoagulation for which there is also less published evidence [14].

Overall, topical hemostatic agents achieve high immediate hemostasis rates (93%, (91%; 94%)). This conclusion is true regardless of the cause of UGIB, including peptic ulcer disease, malignancy, post-endoscopic intervention, and even variceal bleeding. These impressive results remain robust whether the agent is used for primary therapy or rescue therapy. Furthermore, these results are comparable to published effectiveness for other conventional endoscopic modalities such as clips and thermocoagulation (98.5% and 94.5%, respectively) [80]. Direct comparisons between topical hemostatic agents and conventional endoscopic modalities were only available in 10 studies (n=797), with topical agents achieving greater immediate hemostasis (OR 3.94 (1.73; 8.96)). This conclusion remained true when including only data derived from RCT (OR 4.01 (1.52; 10.60). This latter subgroup analysis yields a very important summary result previously not reported by other systematic reviews and for which the small number of patients is counterbalanced by the study quality and magnitude of treatment effect [10-12]. When different topical hemostatic agents were analyzed separately, all studied compounds appeared effective, although the number of patients studied varied greatly.

The major concern about the use of topical hemostatic agents expressed in previous studies has related to the risk of rebleeding. We found that the overall risk of rebleeding was 18% (15%; 21%) among non-comparative studies. This risk is

higher compared to conventional endoscopic approaches [80, 81]. This elevated risk likely represents the inclusion of difficult-to-treat lesions (e.g. large ulcers or difficult position) in which these agents are best used as temporizing strategy until a more definitive endoscopic therapy can be employed. It may also relate to the mechanism of action of some of these agents that form a mechanical barrier to stop bleeding but washes off while a risk of rebleeding may persist (within 24 hours as in the case of Tc-325 and peptic ulcers for which the high-risk period post-hemostasis extends to at least 72 hours [29]). However, studies that directly compared topical agents with conventional endoscopic modalities failed to show a difference in overall rebleeding between the two interventions (OR 1.06 [0.65; 1.74]). Similar conclusions were reached when RCT data only were included, even though the point estimate of overall rebleeding was actually greater for the topical agents, but with a large confidence interval owing to small patient numbers (1.27 [0.62; 2.59]). This observation is congruent with the hypothesis that the risk of rebleeding when using topical agents may have been over-estimated from previous studies, possibly reflecting selection and/or reporting bias. However, it is important to once again emphasize that RCTs comparing topical agents to conventional endoscopic modalities with regards to assessing rebleeding are few, with many exhibiting limitations. For NVU-GIB, there were six RCTs with varying sample sizes. The main limitations with all these studies are two-fold: First, Kwek et al. [39] included a large proportion (60%) of patients with nonbleeding lesions (Forrest IIa and IIb) - lesions to which Hemospray cannot adhere and thus are not indicated for its use. Second, the remaining studies (Baracat et al. [24], Chen et al. [29], Martins et al. [42] and Bang et al. [77]) were small (20-72 patients in each study) and included a large proportion of patients with oozing stigmata (Forrest Ib). This category of lesions has recently been shown to exhibit a lower risk of re-bleeding than previously thought, possibly reflecting the inclusion of lower risk lesions than "true" Forrest Ib [82]. By far, the largest RCT

Study	Interver	ntion	Tota	al	Proportion	95%-Cl	Study	Interventio	on Tota	I	Proportion	95%-Cl
Settina: Hen	nosprav						Settina: En	doClot				
Chen. 2020		7	8		• 0.88	[0.47: 1.00]	Paoluzi, 202	1 1	9		0.11	[0.00: 0.48]
Hussein, 2020	) (3)	0	9.		0.00	0.00: 0.34]	Kim. 2018	2	12	i	0.17	[0.02: 0.48]
Kwek. 2017		3	9		- 0.33	0.07: 0.70]	Hagel. 2020	) 1	18		0.06	[0.00: 0.27]
Pittavanon. 2	016	1	10 -		0.10	0.00: 0.45]	Beg. 2015	1	21	• <u>+</u>	0.05	[0.00: 0.24]
Weaver, 2019	)	0	12 •		0.00	0.00: 0.26]	Vitali, 2019	4	25		0.16	[0.05: 0.36]
Masci, 2014		2	13	_	0.15	0.02: 0.45	Park. 2018	2	37		0.05	[0.01: 0.18]
Arena, 2016		3	15	_	0.20	[0.01: 0.48]	Prei. 2016	6	58		0.10	[0.04: 0.21]
Sulz. 2014		2	15		0.13	[0.02: 0.40]	Subtotal		180	•	0.10	[0.06: 0.16]
Holster, 2012		5	16	_ <u>_</u>	0.31	[0.11: 0.59]	Random eff	fects model		•	0.10	0.06: 0.16
Leblanc, 2013	3	2	17	-	0.12	0.01: 0.36]	Heterogene	ity: I <sup>2</sup> = 0%. P	= 0.76			[]
Malik. 2015		5	17		0.29	0.10: 0.56]	, and a grad			1		
Yau. 2014		7	18	¦	- 0.39	[0.17: 0.64]	Settina: Pu	rastat				
Baracat, 2019	)	5	19		0.26	0.09: 0.51]	Labianca, 20	)21 0	15	• • • • • • • • • • • • • • • • • • •	0.00	[0.00: 0.22]
Disney, 2015		2	19		0.11	[0.01: 0.33]	de Nucci. 20	020 7	41		0.17	[0.07: 0.32]
Sinha, 2016		3	19	_	0.16	[0.03: 0.40]	Subramania	m. 2019 1	44	•   i	0.02	[0.00: 0.12]
Suna, 2011		2	19		0.11	[0.01; 0.33]	Subtotal	, 2015	100		0.12	[0.06: 0.22]
Nasr. 2015		2	21		0.10	[0.01; 0.30]	Random eff	fects model			0.07	[0.01: 0.26]
Meng. 2019		4	25	_	0.16	[0.05: 0.36]	Heterogene	$itv: I^2 = 60\% H$	P = 0.08			[0101, 0120]
Martins 2019	) 1		28		- 0.54	[0.34: 0.72]	incereigene		0.00	i i		
Bestari, 2020		0	30	<u> </u>	0.00	[0.00; 0.12]	Settina: Ne	xpowder		1		
Ibrahim 201	5	1	30 -	•	0.03	[0.00; 0.17]	Park. 2019 (	1) 3	16		0.19	[0.04: 0.46]
Appleby 201	7 1		32	<u> </u>	0.34	[0.19: 0.53]	Shin, 2021	3	40		0.07	[0.02; 0.20]
Paoluzi 2021	. 1	10	32	<b></b>	0.31	[0.16: 0.50]	Park, 2019 (	2) 2	54	i	0.04	[0.00; 0.13]
Hagel, 2017	1	12	33	-	0.36	[0.20: 0.55]	Subtotal		110	🔶 l	0.09	[0.07: 0.16]
Giles. 2016		4	36		0.11	[0.03; 0.26]	Random eff	fects model		-	0.08	[0.03: 0.20]
Widlak, 2015		4	42	-	0.10	[0.03: 0.23]	Heterogene	ity: 1 <sup>2</sup> = 44%. <i>F</i>	P = 0.17		0.00	[0:00, 0:20]
Ibrahim, 2019	9	5	43		0.12	[0.04: 0.25]	··· g-··-	,,.		1		
Thavalasekara	an. 2017	7	44	-	0.16	[0.07: 0.30]	Setting: CE	GP-003				
Gupta, 2018	,	6	45		0.13	0.05: 0.27]	Bang, 2018	3	35		0.09	[0.02: 0.23]
Shivaii. 2018		6	45		0.13	0.05: 0.331						[,]
Prentice, 201	8	9	48		0.19	0.09; 0.33	Total		2696	•	0.22	[0.20; 0.24]
Min, 2018		8	48		0.17	0.07; 0.30]	Random eff	fects model		<b>↓</b>	0.18	0.15: 0.21
Cahyadi, 201	7 2	22	51		0.43	0.29; 0.58]	Prediction i	interval				0.06: 0.43
Smith, 2014		9	55		0.16	0.08; 0.29]	Heterogene	ity: I <sup>2</sup> = 69%, <i>F</i>	<sup>o</sup> <0.01	i.		
Chen, 2015	1	15	56		0.27	0.16; 0.40]	5	,				
Hussein, 2020	0	2	57 ·	•   l	0.04	0.00; 0.12]				0 0.2 0.4 0	0.6 0.8	
Chahal, 2020	2	23	65		0.35	0.24; 0.48]						
Pittayanon, 2	018 2	24	77	-	0.31	0.21; 0.43	Test for sub	group differer	nces (fix	ed effect): $\chi^2_4$	= 29.49, df =	4 (P < 0.01)
Ramirez-polo	,2019 1	12	81		0.15	0.08; 0.24]	Test for sub	group differer	nces (ra	ndom effects)	:	· · ·
Hussein, 202	1(1) 1	13	87	- <b></b>	0.51	0.08; 0.24]	$\chi^2_4 = 14.476$	df = 4 (P < 0.0)	01)	,		
Vitali, 2019	2	24 1	02		0.24	0.16; 0.33	/14	•	,			
Lau, 2021		9 1	58	-	0.08	0.04; 0.15						
Becg, 2020	5	59 1	52		0.39	0.31; 0.47]						
Hussein, 202	1(2) 2	26 1	54		0.17	0.11; 0.24]						
Haddara, 201	6 5	51 1	91		0.27	[0.21; 0.34]						
de Santiago, J	2019 3	35 2	19	<b>.</b>	0.16	[0.11; 0.22]						
Subtotal		22	71	*	0.23 [	0.21; 0.25						
Random effe	cts mode	el		+	0.20	0.17; 0.24]						
Heterogeneit	y: I <sup>2</sup> = 70%	%, P <(	0.01									
			[									
			0	0.2 0.4 0	.6 0.8							

Fig.3 Forrest plot of primary outcome by proportion: overall rebleeding.

assessing Hemospray, was recently published by Lau et al. [15] This study randomized 224 patients with NVUGIB to Hemospray or conventional endoscopic therapy. It concluded that Hemospray was non-inferior to conventional endoscopic hemostasis. However, the study was limited by the inclusion of a large proportion of Forrest lb lesions. Furthermore, this study exhibited a marked imbalance in the malignant bleeding subgroup which were over-represented in the Hemospray arm and may have biased the results in favor of the topical agent [16]. Only one study assessed variceal bleeding (Ibrahim et al. [38]) limiting the conclusions specifically addressing this subgroup of patients.

When the risk of rebleeding was stratified according to the agent used, it was numerically greater with Hemospray (20% [17%; 24%]) compared to other agents (EndoClot 10% [6%; 16%], PuraStat 7% [1%; 27%], or Nexpowder 8% [3%; 20%]) and CEGP-003 (9% [3%; 23%]). This difference may not be statistically significant (there are very few direct comparisons) and may only reflect low numbers of patients treated with the other agents or could alternately be a result of the inclusion of higher-risk lesions in the Hemospray studies. Indeed, only one study

**Table 4** Primary and secondary outcome for comparative studies (comparing topical agent vs. conventional endoscopic therapy).

	No. studies	No.patients	Odds ratio (95 % CI)	<i>P</i> value for heterogeneity	<b> </b> <sup>2</sup>
Primary outcome					
Immediate hemostasis (overall UGIB)	10	797	3.94 (1.73; 8.96)	0.79	0 %
Overall rebleeding	10	775	1.06 (0.65; 1.74)	0.58	0%
Rebleeding 7 days	6	356	0.97 (0.43; 2.16)	0.51	0%
Rebleeding 30 days	7	649	0.75 (0.39; 1.45)	0.24	25%
Secondary outcome (UGIB)					
Overall mortality	9	621	1.05 (0.64; 1.70)	0.79	0%
Bleeding-related mortality	5	347	0.62 (0.14; 2.71)	0.45	0%
Technical success	9 <sup>1</sup>	573	0.30 (0.01; 8.33)	-	-
Adverse events	5 <sup>2</sup>	185	-	-	-

<sup>1</sup> Only one study with estimable data.

<sup>2</sup> All were double zero-event.

	Experir	nental	Cont	rol			
Study	Events	Total	Events	Total	Odds Ratio	OR	95%-Cl
Intervention = Hemospra	у						
Kwek, 2017	9	10	10	10		0.30	[0.01; 8.33]
Sinha, 2016	19	20	16	20		4.75	[0.48; 46.91]
Pittayanon, 2016	10	10	10	10			
Baracat, 2019	19	19	18	20		5.27	[0.24; 117.26]
Chen, 2020	7	8	4	9		8.75	[0.74; 103.82]
Martins, 2019	28	28	31	31			
Lau, 2016	108	111	102	113		3.88	[1.05; 14.32]
Subtotal		206		213		3.62	[1.50; 8.78]
Random effects model						3.78	[1.48; 9.67]
Heterogeneity: $I^2 = 0\%$ , $P =$	0.60						
Intervention = EndoClot							
Beg, 2015	21	21	109	109			
Park, 2018	36	37	128	139		3.09	[0.39; 24.77]
Intervention = CEGP-003							
Bang, 2018	35	35	33	37		9.54	[0.49; 183.98]
Total Random effects model Prediction interval		299		498		3.87 3.94	[1.76; 8.49] [1.73; 8.96] [1.34; 11.57]
Heterogeneity: $I^2 = 0\%$ , $P =$	0.79				0.0 0.01 1 10 100		
Test for subgroup difference Test for subgroup difference	ces (fixed ces (rando	effect): om effec	$\chi_2^2 = 0.42$ ts): $\chi_2^2 = 0$	, df = 2 0.40, d	(P = 0.81) f = 2 (P = 0.82)		

**Fig.4** Forrest plot of primary outcome for comparative studies: immediate hemostasis.

directly compared different topical hemostatic agents (Hemospray vs. EndoClot) [77], concluding similar effectiveness, as well as rebleeding rates and safety profiles when used for UGIB of different etiologies in 127 patients.

When stratified by etiology, the risk of rebleeding was similar in NVUGIB, malignancy-related lesions, as well as in variceal bleeding (Appendix 2b). Even though the risk of rebleeding was numerically greater when the topical agent was used for rescue (25% [20%; 30%]) compared to primary therapy (18%

	Experimental		Control				
Study	Events	Total	Events	Total	Odds Ratio	OR	95%-CI
Intervention = Hemospra	y						
Kwek, 2017	3	9	1	10		4.50	[0.37; 54.16]
Sinha, 2016	3	19	4	16		0.56	[0.11; 3.00]
Pittayanon, 2016	1	10	3	10		0.26	[0.02; 3.06]
Baracat, 2019	5	19	3	18		1.79	[0.36; 8.90]
Chen, 2020	7	8	7	9		2.00	[0.15; 27.45]
Martins, 2019	15	28	12	31		1.83	[0.65; 5.15]
Lau, 2016	9	108	10	102		0.84	[0.33; 2.15]
Subtotal		201		196	-	1.16	[0.68; 1.99]
Random effects model						1.17	[0.67; 2.03]
Heterogeneity: $I^2 = 0\%$ , $P =$	• 0.55						
Intervention = EndoClot							
Beg, 2015	1	21	10	109		0.50	[0.06; 4.09]
Park, 2018	2	37	15	139		0.47	[0.10; 2.16]
Subtotal		58		248		0.48	[0.14; 1.65]
Random effects model						0.48	[0.14; 1.65]
Heterogeneity: I <sup>2</sup> = 0%, <i>P</i> = 0.97							
Intervention = CEGP-003							
Bang, 2018	3	35	1	37		3.38	[0.33; 34.09]
Total		294		481		1.04	[0.65; 1.66]
Random effects model						1.06	[0.65; 1.74]
Prediction interval							[0.59: 1.90]
Heterogeneity: $I^2 = 0\%$ , $P =$	• 0.58						1
					0.0 0.1 0.3 1 2 10		
Test for subgroup differences (fixed effect): $\chi_2^2 = 2.66$ , df = 2 ( $P = 0.26$ )							
lest for subgroup differences (random effects): $\chi_2^2 = 2.66$ , df = 2 ( <i>P</i> = 0.26)							

**Fig.5** Forrest plot of primary outcome for comparative studies: overall rebleeding.

[14%; 22%]); this difference is likely to be non-significant as suggested by marked overlapping confidence intervals. Given the ease of use of these topical agents, they comprise an attractive option to manage UGIB in emergency situations. These compounds may also prove to be the preferred endoscopic modalities when managing diffuse bleeding from fragile surfaces such as in malignancy, especially considering the difficulty and limited success when using conventional endoscopic options in such a scenario [1]. This is supported by the current meta-analysis results as we showed that topical agents were more effective in achieving immediate hemostasis than conventional endoscopic therapy when used for malignancy-related bleeding (OR = 14.74 [2.16;100.61]). This, too, is an important finding that bears important clinical implications and that has not been clearly identified previously using summary data.

One of the major strengths of the topical hemostatic agents is their remarkable safety profile. Indeed, the overall AE rate seen with these agents was 2% (1;3%). Some of the reported AEs were serious but remain rare, such as the three cases of perforations among 2111 patients. Another important advantage when using these agents is the high technical success rate noted (97%, 97;98%). The main technical problem reported was catheter blockage due to premature activation of the powder inside the delivery catheter. This problem was encountered with Hemospray only, and can be avoided by ensuring the endoscope channel and actual delivery catheter are cleared of any liquid (e.g. blood or fluid) before inserting the catheter.

Our study has a number of strengths. It is the largest and most comprehensive meta-analysis performed to-date assessing the effectiveness and safety of different topical hemostatic agents in the management of UGIB. It included 59 studies from different regions of the world published over the past decade, and assessed different agents, hence increasing generalizability. In addition, we assessed the effectiveness of these agents in UGIB of different etiologies (e.g. peptic ulcer disease, malignancy, variceal bleed) and in different settings (primary vs. rescue therapy), further increasing external validity. The main limitation of our analysis is the overall low to very low certainty of the evidence but such limitation exists also with many endoscopic therapeutic modalities yet have not deterred consideration for use in authoritative guidelines with adequate characterization of the data [14]. In addition to limitations in study design, the studies also display a lack of follow-up information in some reports which may have introduced reporting bias. Nevertheless, this systematic review addresses many of the limitations of previously published works that included small number of studies, limited inclusion criteria (limiting generalizability), and no formal assessment of the quality of the evidence using validated instruments such as the GRADE assessment tool [10–12].

#### Conclusions

In conclusion, topical hemostatic agents are effective and safe in managing UGIB of different etiologies even when used as primary therapy and monotherapy. Patients treated with these agents should be monitored closely in the first few days due to a higher risk of rebleeding. More complete and novel subgroup analyses suggest these agents yield the highest immediate hemostatic rates, and are particularly effective in malignant bleeding, even when compared to other modalities. Future well designed studies should further compare hemostatic topical agents to conventional endoscopic modalities, and among each other to better inform recommendations as to their use, especially when chosen as sole first-line agent.

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#### **Competing interests**

The authors declare that they have no conflict of interest.

#### References

- Lau LHS, Sung JJY. Treatment of upper gastrointestinal bleeding in 2020: New techniques and outcomes. Dig Endosc 2021; 33: 83–94
- [2] Abougergi MS, Travis AC, Saltzman JR. The in-hospital mortality rate for upper GI hemorrhage has decreased over 2 decades in the United States: a nationwide analysis. Gastrointest Endosc 2015; 81: 882–888
- [3] Rotondano G. Epidemiology and diagnosis of acute nonvariceal upper gastrointestinal bleeding. Gastroenterol Clin North Am 2014; 43: 643–663
- [4] Lau JY, Barkun A, Fan DM et al. Challenges in the management of acute peptic ulcer bleeding. Lancet 2013; 381: 2033–2043
- [5] Hearnshaw SA, Logan RF, Lowe D et al. Acute upper gastrointestinal bleeding in the UK: patient characteristics, diagnoses and outcomes in the 2007 UK audit. Gut 2011; 60: 1327–1335
- [6] Barkun AN, Moosavi S, Martel M. Topical hemostatic agents: a systematic review with particular emphasis on endoscopic application in Gl bleeding. Gastrointest Endosc 2013; 77: 692–700
- [7] Chen YI, Barkun AN. Hemostatic powders in gastrointestinal bleeding: A systematic review. Gastrointest Endosc Clin N Am 2015; 25: 535–552
- [8] Pioche M, Camus M, Rivory J et al. A self-assembling matrix-forming gel can be easily and safely applied to prevent delayed bleeding after endoscopic resections. Endosc Int Open 2016; 4: E415–E419
- [9] de Rezende DT, Brunaldi VO, Bernardo WM et al. Use of hemostatic powder in treatment of upper gastrointestinal bleeding: a systematic review and meta-analysis. Endosc Int Open 2019; 7: E1704–E1713

- [10] Mutneja H, Bhurwal A, Go A et al. Efficacy of Hemospray in upper gastrointestinal bleeding: A systematic review and meta-analysis. J Gastrointestin Liver Dis 2020; 29: 69–76
- [11] Aziz M, Weissman S, Mehta TI et al. Efficacy of Hemospray in nonvariceal upper gastrointestinal bleeding: A systematic review with meta-analysis. Ann Gastroenterol 2020; 33: 145–154
- [12] Facciorusso A, Straus TM, Eyileten PC et al. Efficacy of hemostatic powders in upper gastrointestinal bleeding: A systematic review and meta-analysis. Dig Liver Dis 2019; 51: 1633–1640
- [13] Barkun AN, Almadi M, Kuipers EJ et al. Management of nonvariceal upper gastrointestinal bleeding: guideline recommendations from the International Consensus Group. Ann Intern Med 2019; 171: 805– 822
- [14] Laine L, Barkun AN, Saltzman JR et al. ACG Clinical Guideline: Upper Gastrointestinal and Ulcer Bleeding. Am J Gastroenterol 2021; 116: 899–917
- [15] Lau JYW, Pittayanon R, Kwek A et al. Comparison of a hemostatic powder and standard treatment in the control of active bleeding from upper nonvariceal lesions: A multicenter, noninferiority, randomized trial. Ann Intern Med 2022; 175: 171–178
- [16] Barkun AN, Alali A. The role of hemostatic powder in endoscopic hemostasis of nonvariceal upper gastrointestinal bleeding. Ann Intern Med 2022; 175: 289–290
- [17] Higgins JP, Altman DG, Gotzsche PC et al. The Cochrane Collaboration's tool for assessing risk of bias in randomised trials. BMJ 2011; 343: d5928
- [18] Wells G, Shea B, O'Connell D et al. The Newcastle Ottawa Scale (NOS) for assessing the quality of nonrandomised studies in metaanalyses. https://www.ohri.ca/programs/clinical\_epidemiology/oxford.asp
- [19] Guyatt GH, Oxman AD, Vist GE et al. GRADE: An emerging consensus on rating quality of evidence and strength of recommendations. BMJ 2008; 336: 924–926
- [20] Laine L, Spiegel B, Rostom A et al. Methodology for randomized trials of patients with nonvariceal upper gastrointestinal bleeding: Recommendations from an international consensus conference. Am J Gastroenterol 2010; 105: 540–550
- [21] Subramaniam S, Kandiah K, Chedgy F et al. A novel self-assembling peptide for hemostasis during endoscopic submucosal dissection: A randomized controlled trial. Endoscopy 2021; 53: 27–35
- [22] Appleby R, Hoare J. Hemospray in a large tertiary NHS trust: A descriptive analysis of the first three years of use. Gut 2017; 66: A216
- [23] Arena M, Masci E, Eusebi LH et al. Hemospray for treatment of acute bleeding due to upper gastrointestinal tumours. Dig Liver Dis 2017; 49: 514–517
- [24] Baracat FI, de Moura DTH, Brunaldi VO et al. Randomized controlled trial of hemostatic powder versus endoscopic clipping for non-variceal upper gastrointestinal bleeding. Surg Endosc 2020; 34: 317–324
- [25] Becq A, Houdeville C, Tran Minh ML et al. Experience with the use of a hemostatic powder in 152 patients undergoing urgent endoscopy for gastrointestinal bleeding. Clin Res Hepatol Gastroenterol 2020; 45: 101558
- [26] Bestari MB, Joewono IR, Girawan D et al. Hemospray during emergency endoscopy: indonesia's first experience from 37 patients. Case Rep Gastroenterol 2020; 14: 70–79
- [27] Cahyadi O, Bauder M, Meier B et al. Effectiveness of TC-325 (Hemospray) for treatment of diffuse or refractory upper gastrointestinal bleeding – a single center experience. Endosc Int Open 2017; 5: E1159–E1164
- [28] Chahal D, Lee JGH, Ali-Mohamad N et al. High rate of re-bleeding after application of Hemospray for upper and lower gastrointestinal bleeds. Dig Liver Dis 2020; 52: 768–772

- [29] Chen YI, Barkun A, Nolan S. Hemostatic powder TC-325 in the management of upper and lower gastrointestinal bleeding: A two-year experience at a single institution. Endoscopy 2015; 47: 167–171
- [30] Chen YI, Wyse J, Lu Y et al. TC-325 hemostatic powder versus current standard of care in managing malignant GI bleeding: A pilot randomized clinical trial. Gastrointest Endosc 2020; 91: 321–328 e1
- [31] Disney B, Kurup A, Muhammad H et al. Hemospray use for the management of acute bleeding from upper gastrointestinal cancer: The Russells Hall experience. Gut 2015; 64: A71–A72
- [32] Giles H, Lal D, Gerred S et al. Efficacy and safety of TC-325 (Hemospray) for non-variceal upper gastrointestinal bleeding at Middlemore Hospital: The early New Zealand experience. N Z Med J 2016; 129: 38– 43
- [33] Gupta T, Leet F, Harborne M et al. Hemospray use in acute gastrointestinal bleeding – a 4 year single centre experience. Gut 2018; 67: A39
- [34] Haddara S, Jacques J, Lecleire S et al. A novel hemostatic powder for upper gastrointestinal bleeding: a multicenter study (the "GRAPHE" registry). Endoscopy 2016; 48: 1084–1095
- [35] Hagel AF, Albrecht H, Nagel A et al. The application of Hemospray in gastrointestinal bleeding during emergency endoscopy. Gastroenterol Res Pract 2017; 2017: 3083481
- [36] Holster IL, Poley JW, Kuipers EJ et al. Controlling gastric variceal bleeding with endoscopically applied hemostatic powder (Hemospray). J Hepatol 2012; 57: 1397–1398
- [37] Ibrahim M, El-Mikkawy A, Abdalla H et al. Management of acute variceal bleeding using hemostatic powder. United European Gastroenterol J 2015; 3: 277–283
- [38] Ibrahim M, El-Mikkawy A, Abdel Hamid M et al. Early application of haemostatic powder added to standard management for oesophagogastric variceal bleeding: A randomised trial. Gut 2019; 68: 844– 853
- [39] Kwek BEA, Ang TL, Ong PLJ et al. TC-325 versus the conventional combined technique for endoscopic treatment of peptic ulcers with high-risk bleeding stigmata: A randomized pilot study. J Dig Dis 2017; 18: 323–329
- [40] Leblanc S, Vienne A, Dhooge M et al. Early experience with a novel hemostatic powder used to treat upper GI bleeding related to malignancies or after therapeutic interventions (with videos). Gastrointest Endosc 2013; 78: 169–175
- [41] Malik A, Duane P, Eadala P. Use of Hemospray for non variceal upper gastrointestinal bleed in a district general hospital. Gut 2015; 64: A64
- [42] Martins B, Corsato R, Batista Pires C et al. Preliminary results of a randomized controlled trial comparing hemostatic powder versus optimal clinical treatment in the management. United Europ Gastroenterol J 2019; 8S: P0135
- [43] Masci E, Arena M, Morandi E et al. Upper gastrointestinal active bleeding ulcers: Review of literature on the results of endoscopic techniques and our experience with Hemospray. Scand J Gastroenterol 2014; 49: 1290–1295
- [44] Meng ZW, Marr KJ, Mohamed R et al. Long-term effectiveness, safety and mortality associated with the use of TC-325 for malignancyrelated upper gastrointestinal bleeds: A multicentre retrospective study. J Can Assoc Gastroenterol 2019; 2: 91–97
- [45] Min T, Mothey M, Sharrack N et al. Efficacy of haemospray therapy on re-bleed and mortality rates – a UK single-centre experience. United Europ Gastroenterol J 2018; 85: P0773
- [46] Nasr I, De Martino S, Borrow D et al. PWE-007 When to use hemospray? A single centre experience Gut 2015; 64 (supp1): A212–A213
- [47] Pittayanon R, Prueksapanich P, Rerknimitr R. The efficacy of Hemospray in patients with upper gastrointestinal bleeding from tumor. Endosc Int Open 2016; 4: E933–E936

- [48] Pittayanon R, Rerknimitr R, Barkun A. Prognostic factors affecting outcomes in patients with malignant GI bleeding treated with a novel endoscopically delivered hemostatic powder. Gastrointest Endosc 2018; 87: 994–1002
- [49] Prentice E, Masegosa Ataz J, Barron M et al. A retrospective singlecentre review of hemospray in GI bleeding. United Europ Gastroenterol J 2018; 8S: P1105
- [50] Ramirez-Polo AI, Casal-Sanchez J, Hernandez-Guerrero A et al. Treatment of gastrointestinal bleeding with hemostatic powder (TC-325): A multicenter study. Surg Endosc 2019; 33: 2349–2356
- [51] Shivaji U, Leet F, Gupta T et al. Hemospray use in acute gastrointestinal bleeding – a 4-year single-centre experience. United Europ Gastroenterol J 2018; 8S: P1726
- [52] Sinha R, Lockman KA, Church NI et al. The use of hemostatic spray as an adjunct to conventional hemostatic measures in high-risk nonvariceal upper GI bleeding (with video). Gastrointest Endosc 2016; 84: 900–906 e3
- [53] Smith LA, Stanley AJ, Bergman JJ et al. Hemospray application in nonvariceal upper gastrointestinal bleeding: Results of the Survey to Evaluate the Application of Hemospray in the Luminal Tract. J Clin Gastroenterol 2014; 48: e89–e92
- [54] Sulz MC, Frei R, Meyenberger C et al. Routine use of Hemospray for gastrointestinal bleeding: Prospective two-center experience in Switzerland. Endoscopy 2014; 46: 619–624
- [55] Sung JJ, Luo D, Wu JC et al. Early clinical experience of the safety and effectiveness of Hemospray in achieving hemostasis in patients with acute peptic ulcer bleeding. Endoscopy 2011; 43: 291–295
- [56] Thayalasekaran S, Bhandari P, Quine A et al. Hemospray use in the management of upper gastrointestinal haemorrhage: A 2-year experience across 2 teaching hospitals in the north and south of England. Gut 2017; 85: AB35
- [57] Weaver M, Richard R, Rood M et al. Clinical outcomes and endoscopist's perception of the use of hemospray for gastrointestinal bleeding: Outcomes from a U.S. academic tertiary care medical center. Am J Gastroenterol 2019; 114 : S363
- [58] Widlak M, Wijesinghe H, Siau K et al. PWE-032 Hemospray for acute upper gastrointestinal bleeding – a single centre experience. Gut 2015; 64 (supp1): A225
- [59] Yau AH, Ou G, Galorport C et al. Safety and efficacy of Hemospray in upper gastrointestinal bleeding. Can J Gastroenterol Hepatol 2014; 28: 72–76
- [60] Rodriguez de Santiago E, Burgos-Santamaria D, Perez-Carazo L et al. Hemostatic spray powder TC-325 for GI bleeding in a nationwide study: Survival and predictors of failure via competing risks analysis. Gastrointest Endosc 2019; 90: 581–590 e6
- [61] Hussein M, Alzoubaidi D, Lopez MF et al. Hemostatic spray powder TC-325 in the primary endoscopic treatment of peptic ulcer-related bleeding: multicenter international registry. Endoscopy 2021; 53: 36–43
- [62] Hussein M, Alzoubaidi D, O'Donnell M et al. Hemostatic powder TC-325 treatment of malignancy-related upper gastrointestinal bleeds: International registry outcomes. J Gastroenterol Hepatol 2021; 36: 3027–3032
- [63] Hussein M, Alzoubaidi D, Serna A et al. Outcomes of Hemospray therapy in the treatment of intraprocedural upper gastrointestinal bleeding post-endoscopic therapy. United Europ Gastroenterol J 2020; 8: 1155–1162
- [64] Hussein M, Alzoubaidi D, Weaver M et al. Hemospray in the treatment of variceal bleeds: outcomes from the international Hemospray registry. Gut 2021; 70: A1–A262
- [65] Beg S, Al-Bakir I, Bhuva M et al. Early clinical experience of the safety and efficacy of EndoClot in the management of non-variceal upper gastrointestinal bleeding. Endosc Int Open 2015; 3: E605–E609

- [66] Kim YJ, Park JC, Kim EH et al. Hemostatic powder application for control of acute upper gastrointestinal bleeding in patients with gastric malignancy. Endosc Int Open 2018; 6: E700–E705
- [67] Prei JC, Barmeyer C, Burgel N et al. EndoClot polysaccharide hemostatic system in nonvariceal gastrointestinal bleeding: Results of a prospective multicenter observational pilot study. J Clin Gastroenterol 2016; 50: e95–e100
- [68] Hagel AF, Raithel M, Hempen P et al. Multicenter analysis of endoclot as hemostatic powder in different endoscopic settings of the upper gastrointestinal tract. J Physiol Pharmacol 2020; 71: 657–664
- [69] Park JC, Kim YJ, Kim EH et al. Effectiveness of the polysaccharide hemostatic powder in non-variceal upper gastrointestinal bleeding: Using propensity score matching. J Gastroenterol Hepatol 2018; 33: 1500–1506
- [70] de Nucci G, Reati R, Arena I et al. Efficacy of a novel self-assembling peptide hemostatic gel as rescue therapy for refractory acute gastrointestinal bleeding. Endoscopy 2020; 52: 773–779
- [71] Subramaniam S, Kandiah K, Thayalasekaran S et al. Haemostasis and prevention of bleeding related to ER: The role of a novel self-assembling peptide. United Europ Gastroenterol J 2019; 7: 155–162
- [72] Branchi F, Klingenberg-Noftz R, Friedrich K et al. PuraStat in gastrointestinal bleeding: results of a prospective multicentre observational pilot study. Surg Endosc 2022; 36: 2954–2961
- [73] Labianca O, Sica M, Zulli C et al. Use of Purastat in management of gastro-intestinal bleeding: our experience. Endoscopy 2021; 53: S97– S98

- [74] Park JS, Bang BW, Hong SJ et al. Efficacy of a novel hemostatic adhesive powder in patients with refractory upper gastrointestinal bleeding: A pilot study. Endoscopy 2019; 51: 458–462
- [75] Park J-S, Lee D-H, Hong S-J et al. Novel hemostatic adhesive powder application in nonvariceal upper gastrointestinal bleeding. UEG J 2019; 7: 133–134
- [76] Shin J, Cha B, Park JS et al. Efficacy of a novel hemostatic adhesive powder in patients with upper gastrointestinal tumor bleeding. BMC Gastroenterol 2021; 21: 40
- [77] Bang BW, Lee DH, Kim HK et al. CEGP-003 spray has a similar hemostatic effect to epinephrine injection in cases of acute upper gastrointestinal bleeding. Dig Dis Sci 2018; 63: 3026–3032
- [78] Vitali F, Naegel A, Atreya R et al. Comparison of Hemospray and Endoclot for the treatment of gastrointestinal bleeding. World J Gastroenterol 2019; 25: 1592–1602
- [79] Paoluzi OA, Cardamone C, Aucello A et al. Efficacy of hemostatic powders as monotherapy or rescue therapy in gastrointestinal bleeding related to neoplastic or non-neoplastic lesions. Scand J Gastroenterol 2021; 56: 1–8
- [80] Sung JJ, Tsoi KK, Lai LH et al. Endoscopic clipping versus injection and thermo-coagulation in the treatment of non-variceal upper gastrointestinal bleeding: A meta-analysis. Gut 2007; 56: 1364–1373
- [81] Bratanic A, Puljiz Z, Ljubicicz N et al. Predictive factors of rebleeding and mortality following endoscopic hemostasis in bleeding peptic ulcers. Hepatogastroenterology 2013; 60: 112–117
- [82] Jensen DM, Eklund S, Persson T et al. Reassessment of rebleeding risk of forrest ib (oozing) peptic ulcer bleeding in a large international randomized trial. Am J Gastroenterol 2017; 112: 441–446