Comparison of dextran and albumin on blood coagulation in patients undergoing major gynaecological surgery

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Abstract

Background: Hydroxyethyl starches have been withdrawn from the European market. In Sweden, dextran was the main colloid until 2000, when starches overtook the market. After the recent 6S-trial, it was suggested that dextran could be reinstituted, but concerns for greater coagulopathy, bleeding and anaphylaxis still remain. An experimental study from our department indicated that isovolemic substitution of dextran-70 did not derange the von Willebrand function more than albumin 5%, considering the fact that dextran is hyperoncotic in comparison to albumin 5% and, therefore, induces a greater plasma volume expansion and thereby a greater dilutional coagulopathy.

Methods: Eighteen patients undergoing major gynaecological surgery were assigned to receive either 5% albumin or 6% dextran-70 with 9 patients in each group. Standard coagulation tests, including prothrombin time (PT), activated partial thromboplastin time (aPTT), fibrinogen and platelet count, viscoelastic coagulation test thromboelastometry (ROTEM) and the Multiplate platelet aggregation test were used to test for coagulation defects at different time points perioperatively. Blood loss, blood loss replacement data and haemodynamic parameters were retrieved from anaesthetic and postoperative charts. A local departmental fluid and transfusion/infusion protocol assured haemoglobin > 90 g/l and mean arterial pressure > 65 mmHg with Ringer’s acetate in addition to the colloid use.

Results: There were no differences in demographic data between the groups. The tissue factor-activated (EXTEM) clot-structure parameter ROTEM A10 was decreased significantly in the dextran group as compared to the albumin group after the infusion of 500 ml of either colloid solution. The PT and aPTT were significantly prolonged, and the platelet count decreased postoperatively in the dextran group, whereas albumin only deranged fibrinogen levels as compared to preoperative levels. There were no differences in Multiplate platelet aggregometry, amount of haemorrhage or transfusion of blood components between the groups.

Conclusions: Standard plasma-based coagulation tests, platelet count and whole blood viscoelastic clot structure are affected by 6% dextran-70 to a greater extent than by 5% albumin, but platelet aggregation is not. Future studies should use more advanced haemodynamic monitoring to assess isovolemic plasma volume expansion with dextran and whether this affects haemostasis to a lesser degree.

Keywords: Dextran, Albumin, Surgery, Blood-loss replacement, Fluid therapy, Gynaecologic cancer

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Background

Patients undergoing major gynaecological cancer resection surgery are at risk for fluid loss through evaporation, bleeding, capillary leakage and ascites formation. Crystalloids are often used for the initial fluid management; then colloids are added upon larger fluid/initial blood losses (Kristensen et al. 2014). The benefits of colloid infusion over crystalloid are still debated. A Cochrane review comparing the use of crystalloids and colloids in fluid resuscitation of critically ill patients did not show any reduction in mortality with the use of colloids. The authors argued that crystalloids should be used since they are less expensive (Perel et al. 2013).

Still, colloids are often used for perioperative fluid therapy due to their better volume-expanding effects and better sustained volume effect with less risk for interstitial oedema affecting different organs (Vercueil et al. 2005). Another Cochrane review that compared the various colloid solutions available on the market showed no advantages of one over the other, in terms of safety and efficacy (Bunn and Trivedi 2012).

Dextran was the leading synthetic colloid in Sweden until 2001, but preference then shifted towards hydroxyethyl starch (HES) (Berseus et al. 2008). The disadvantages of dextran on haemostasis, an increased risk for anaphylaxis as compared to other synthetic colloids and the switch to low molecular weight heparins for thromboprophylaxis also contributed to this shift from dextran to starch.

There are only two recent randomised clinical studies on haemostasis and bleeding complications after perioperative dextran use (Rasmussen et al. 2015; Zdolsek et al. 2011).

A retrospective study on sepsis patients from Copenhagen, Denmark, showed greater bleeding (Hvidt and Perner 2012), while an older study on trauma patients did not indicate greater bleeding with the use of dextran (Modig 1986). A recent experimental study from our institution that focussed on comparing albumin with dextran during the isovolemic blood volume restitution of blood losses did not find any major differences in thromboelastometry (ROTEM)® or von Willebrand factor analyses (Schött et al. 2017).

In a survey study, HES was the most used colloid in intensive care units in Scandinavia in 2008 (Perner et al. 2008). With growing concern regarding the safety of HES and its recent suspension from the market (Agency 2018), an increased use of albumin has been noted throughout Scandinavia (Kongsgaard et al. 2018). Albumin is relatively expensive but has shown increased survival in septic patients in a recent meta-analysis (Delaney et al. 2011). However, albumin has also been shown to impair coagulation and platelet function (De Jonge and Levi 2001; Li et al. 2015). Dextran has been used at our hospital, largely due to local routines and ongoing research in our experimental laboratory (Schött et al. 2017; Persson and Grände 2006; Dubniks et al. 2009).

The aim of this study was to see if there was any difference between fluid regimes with 6% dextran-70 or 5% albumin in major gynaecological surgery regarding routine coagulation tests, viscoelastic coagulation test ROTEM®, platelet function measured with a new bedside platelet aggregometer Multiplate®, bleeding or need for transfusion. We focused on patients undergoing surgery for ovarian cancer since this type of surgery is very standardised at our hospital, which leads to more comparable groups. They have advanced fluid shifts and are often hypovolemic, which is addressed in a recently finished study on albumin infusion rates to correct plasma volume deficits (Statkevicius et al. 2016). Also, since these patients often are hypercoagulable, surgeons were more positive to us, using dextran in comparison to albumin. Hypercoagulable patients probably have a better dilutive capacity before a colloid induced coagulopathy could endanger perioperative haemostasis (Nilsson et al. 2016; Thomas et al. 2016).

Methods

Ethical approval was obtained from the regional ethical review board (Lund, Protocol DNR 2010/482). Eighteen patients admitted to the University Hospital in Lund for elective gynaecological cancer operation between February 2014 and April 2014 were included in the study. All the patients’ informed and signed consent were retrieved before the operation.

The patients were anaesthetised in a standardised fashion. Fentanyl, propofol and rocuronium were used for induction, and sevoflurane was used to uphold anaesthesia. In cases where epidural analgesia was not used or was insufficient, fentanyl was given intraoperatively for pain management, and morphine was used for patient-controlled anaesthesia postoperatively.

Glucose 2.5% was used at a background infusion rate of 1 ml/kg/h intraoperatively and in the recovery ward as a routine in our hospital and is a part of the Air study protocol addressing the same type of patients (see above) (Statkevicius et al. 2016). Ringer’s acetate was infused at a basal rate of 4 ml/kg/h and could be increased if deemed necessary to maintain hourly diuresis at > 0.5 ml/kg of bodyweight and mean arterial pressure (MAP) of > 65 mmHg.

A local departmental fluid and transfusion protocol with packed red cells (PRBC) assured a haemoglobin > 90 g/l.

The colloid fluid was chosen by patient order—the first patient received albumin, the second dextran, the third albumin, and so on after consent approval from...
the patient and inclusion criteria were met. Either 5% albumin (CSL Behring, Germany) or 6% dextran (Macrodex®, Meda, Sweden) were used. Albumin was infused in consecutive batches of 250 ml and dextran in 500 ml batches. Dextran was restricted to 1000 ml to ensure patient haemostatic safety, as in vitro studies have indicated effects on coagulation and platelet function after only 500 ml (Aberg et al. 1979). Daily dextran infusions should be restricted to 500–1500 ml or 10–30 ml/kg of body weight (BW), according to the Pharmaceutical Specialties in Sweden (FASS). The FASS does not restrict albumin dosage according to haemostatic safety parameters, but it is well known that a dilutive coagulopathy and platelet dysfunction can be induced (De Jonge and Levi 2001). In vitro, this can be seen at 30% dilution (Nilsson et al. 2016), corresponding to a 1250 ml albumin in vivo infusion (calculated from female blood volume of 60 ml/kg BW and a median weight of 65 kg = 4000 ml; 30% dilution = 1200 ml). Therefore, we restricted the albumin dose to 1250 ml in this study.

The patients’ blood was sampled preoperatively (pre surgery), direct postoperatively (post surgery) and on the day following surgery (day 1). In addition, the blood was sampled after 500 ml colloid and 1000 ml colloid for ROTEM and Multiplate analyses.

The blood samples were drawn from an indwelling radial arterial catheter with a continuous flush and with a sampling membrane, which eliminated the need for disposing blood samples before sampling with the vacutainer technique.

The samples were analysed at each time point by conventional coagulation tests: prothrombin time (PT), activated partial thromboplastin time (aPTT), platelet count and fibrinogen, along with ROTEM and Multiplate. The blood samples for the ROTEM, aPTT, PT and fibrinogen tests were collected in 2 × 2.7 ml tubes containing 0.109 M citrate (BD, UK). The blood for the platelet counts was sampled in 3.0 ml EDTA-tubes. The blood for the Multiplate analyses was collected in 3.0 ml tubes containing recombinant hirudin (Dynabyte GmbH, Munich, Germany).

The PT, aPTT, fibrinogen and platelet count analyses were performed at an accredited hospital laboratory. The PT test was performed using a combined thromboplastin reagent (Stago prothrombin complex assay, SPA +, Stago). The Owren PT assay was calibrated using international normalised ratio (INR) calibrators certified by the Swedish external quality assessment organisation (Equalis, Uppsala, Sweden). The reference range for PT-INR is 0.9–1.2.

The aPTT was analysed with an aPTT reagent from Actin FSL (Siemens Healthcare Diagnostics). The reference range for aPTT has been established locally to 26–33 s. The plasma fibrinogen concentration was measured using the Dade Thrombin reagent (Siemens Healthcare Diagnostics, CS-5100). The local reference range for fibrinogen is set to 2–4 g/l.

The platelet counts were measured using the Sysmex XE 5000 cell counter (Sysmex Corp., Kobe, Japan). The locally determined reference range for platelet count is 165–387 × 10^9/l for adult women.

The thromboelastometry was carried out using the rotational thromboelastometry (ROTEM®; TEM Innovations GmbH, Germany) instrument according to the manufacturer’s instructions. The ROTEM assays were run for 60 min, and samples were analysed with two different ROTEM tests: EXTEM and FIBTEM. The variables that were registered for EXTEM (normal range within brackets) were CT/clotting time (42–74 s), CFT/clot formation time (46–148 s), AA/alpha-angle (63–81°), A10/clot formation at 10 min (43–65 mm) and MCF/maximum clot formation (63–81 mm). For FIBTEM, A10 (9–24 mm) and MCF (9–25 mm) were analysed.

The impedance aggregometry was carried out using the Multiplate® analyser (Roche Diagnostics, Basel, Switzerland) according to the manufacturer’s instructions. Three platelet receptor agonists were used: adenosine phosphate (ADP), collagen (COL) and a thrombin receptor activating peptide (TRAP), and area under the curve (AUC) were measured (normal range within brackets): ADP-test (53–122), COL-test (46–117) and TRAP-test (94–156).

The statistical analyses were performed using Graphpad Prism 6. The Wilcoxon rank-sum test was used to calculate the changes in coagulation parameters by comparing the post 500 ml colloids, post 1000 ml colloids and post surgery and day 1 values to the control pre surgery value (only for dextran due to inadequate patient numbers receiving 1000 ml albumin infusions; n = 2). The Mann-Whitney U test was used to calculate the changes between dextran and albumin groups. Due to multiple statistical calculations, the p-values are presented after Bonferroni correction. p < 0.05 was considered significant. The median values are given, and the results are presented as tables and boxplots. The data was not regarded as normal distributed.

**Results**

Eighteen patients were included in this study. All had ovarian cancer in advanced stage; therefore, surgery involved ovarian resection, hysterectomy and lymph node removals. Seven patients received dextran, and seven received albumin. Four patients did not require volume support with colloids. Calculating statistical power from 14 to 18 patients and dilutive effects on fibrinogen and differences between albumin and dextran on ROTEM A10 indicates that our pilot study is underpowered at a 0.65–0.75 level. There were no significant differences in
age, bleeding or duration of surgery; amount of fluids,
colloids or units (250 ml) of PRBC given; or numbers of
postoperative complications (Table 1).

The ROTEM EXTEM intragroup changes for both
groups are presented in Fig. 1. There were no significant
changes ($p < 0.05$). For the dextran group, the CFT in-
creased from 51 s (s) pre surgery to 86 s after infusion
of 500 ml of dextran and increased to 89 s post surgery.
The alpha-angle decreased from 80° pre surgery to 73°
after infusion of 500 ml of dextran. The EXTEM A10
decreased from a pre-surgery value of 69 mm (mm) to
54 mm after infusion of 500 ml of dextran. The MCF
decreased from 74 mm pre surgery to 65 mm after
infusion of 500 ml of dextran. For the albumin group,
the CFT increased from 72 s pre surgery to 92 s post
surgery. The A10 fell from 62 s pre surgery to 56 s post
surgery.

The FIBTEM A10 and MCF intragroup changes for
dextran and albumin are also presented in Fig. 1. There
were no significant changes ($p < 0.05$). For the dextran
group, the FIBTEM A10 fell from 24 mm pre surgery to
10 mm after infusion of 500 ml of dextran and to 9 mm
post surgery. The MCF fell from 25 mm pre surgery to
12 mm after infusion of 500 ml of dextran and to
10 mm post surgery. For the albumin group, the A10 fell
from 17 mm pre surgery to 11 mm after infusion of
500 ml of albumin. The MCF fell from 19 mm pre sur-
gery to 12 mm after infusion of 500 ml of albumin.

The changes over time for the two groups, expressed
as Δ values for A10 and MCF (EXTEM + FIBTEM), are
presented in Table 2. A significant intergroup difference
($; p < 0.05$) between pre surgery and after the infusion
of 500 ml of dextran was seen for EXTEM A10 ($p =
0.0456$), and the difference remained almost significant
until post surgery ($p = 0.0594$).

There were no significant differences ($p < 0.05$) in the
Multiplate AUC values for either of the groups (Table 3).

The changes in routine coagulation values for both
groups are presented in Table 4. In the dextran group, the
platelet count significantly decreased from $354 \times 10^9/\text{l}$ pre

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### Table 1

| Demographics of the two different groups. No significant differences ($p > 0.05$) between the dextran and albumin substitution groups. Packed red blood cells (PRBCs) |
|------------------|------------------|------------------|
|                  | Dextran $n = 7$  | Albumin $n = 7$  |
| Age (Average)    | 61               | 62.9             |
| Age (Median)     | 59               | 65               |
| Age (Min-Max)    | (54–78)          | (45–77)          |
| Bleeding during surgery (ml) (Average) | 1110             | 679              |
| Bleeding during surgery (ml) (Median) | 900              | 750              |
| Bleeding during surgery (ml) (Min-Max) | (200–3500)       | (200–1100)       |
| Surgery time (minutes) (Average) | 281              | 325              |
| Surgery time (minutes) (Median) | 305              | 358              |
| Surgery time (minutes) (Min-Max) | (162–369)        | (133–411)        |
| Total amount of fluid given (ml) (Average) | 4350             | 5040             |
| Total amount of fluid given (ml) (Median) | 4580             | 4270             |
| Total amount of fluid given (ml) (Min-Max) | (2400–6950)      | (2260–8940)      |
| Total amount of colloids given (ml) (Average) | 982              | 821              |
| Total amount of colloids given (ml) (Median) | 1000             | 750              |
| Total amount of colloids given (ml) (Min-Max) | (500–1750)       | (250–1250)       |
| Total amount of PRBCs given (units) (Average) | 1.57             | 0.43             |
| Total amount of PRBCs given (units) (Median) | 0                | 0                |
| Total amount of PRBCs given (units) (Min-Max) | (0–6)            | (0–2)            |
| Number of patients with postoperative complications | 5               | 6                |
surgery to $317 \times 10^9/l$ ($p = 0.0312$) post surgery and to $221 \times 10^9/l$ ($p = 0.0312$) on day one. The PT-INR increased from 1.0 pre surgery to 1.2 ($p = 0.0312$) post surgery to 1.3 ($p = 0.0312$) on day one, and the aPTT changed from 25 s pre surgery to 31 s ($p = 0.0312$) post surgery and to 30 s ($p = 0.0312$) on day one. The fibrinogen changed from 4.2 g/l pre surgery to 1.9 g/l ($p = 0.0312$) post surgery.

In the albumin group, the platelet count decreased from $328 \times 10^9/l$ pre surgery to $118 \times 10^9/l$ ($p = 0.0312$) on day one. The PT-INR was prolonged from 1.0 pre surgery to 1.3 ($p = 0.0312$) on day one. The fibrinogen decreased from 4.0 g/l pre surgery to 2.1 g/l ($p = 0.0312$) post surgery.

**Discussion**

Signs of impaired coagulation were seen already after infusion of 500 ml of dextran when measured with ROTEM and traditional coagulation tests. Statistically significant changes in the routine laboratory values were observed in both groups. The dextran group showed significant intragroup decreases in both platelet counts and fibrinogen levels, as well as in the PT-INR and aPTT, measured pre surgery, post surgery and on day one after surgery.

The albumin group showed similar tendencies, though first statistically significant on day one post surgery. There were no significant differences between the groups in bleeding, amount of administered crystalloids, colloids or blood products or numbers of postoperative complications. When valuing the changes over time, presented as $\Delta$ values and illustrated in Table 2, a significant decrease in EXTEM A10 was observed after 500 ml infusion in the dextran group, whereas the decrease in
and plasma volume expanding effect. Compared to HES, dextran probably related to its longer half-time dured longer postoperatively (not significantly lower as described in Table 4, with normalised fibrinogen levels at day 1.)

The measurements illustrated in Fig. 1 show a decrease in the FIBTEM A10 and MCF in both groups. Two patients in the dextran group and one patient in the albumin group had values of A10 < 7 mm and MCF < 10 mm, which are often regarded as trigger values for corrective treatment with plasma or fibrinogen concentrate in transfusion protocols using ROTEM (Bolliger et al. 2012). Fibrinogen concentrate appears to reverse the FIBTEM defects more potently after albumin dilution than after dextran dilution in vitro (Winstedt et al. 2013). No patients in our study received fibrinogen concentrate. Being an acute phase reactant, fibrinogen plasma levels start to recover immediately after bleeding and colloid infusions have stopped (Nilsson et al. 2016; Thomas et al. 2016; Wyatt et al. 2013). This can be seen in Table 4, with normalised fibrinogen levels at day 1.

The recent trend shift in colloid consumption following a dramatic reduction in HES sales has brought other solutions back into the limelight. Albumin use has increased steadily in the Nordic countries since 2012, while dextran has seen sales drop by more than 40% over the same period (Kongsgaard et al. 2018).

As mentioned earlier, dextran is known to impair coagulation through various mechanisms, with a risk of bleeding. Dextran induces the so-called von Willebrand-syndrome, which signifies a decrease in the levels/activity of the von Willebrand factor (vWF) as well as factor VIII (Aberg et al. 1979). Fibrin polymerisation is complex and other factors than colloids, like smoking, genetics, tumour induced hypercoaguability, atherosclerosis and medications can be involved (Weisel and Litvinov 2013).
The mechanism behind this phenomenon is not fully understood and has even been observed with other artificial colloids. One theory is that macromolecules present in colloids bind to vWF, thus, accelerating its elimination (Treib et al. 1997; de Jonge et al. 1998). Additionally, dextran affects fibrinogen polymerisation by co-binding, increases the circulating levels of tissue type plasminogen activator (t-PA) and decreases the levels of fibrinolysis plasminogen activator inhibitor-1 (PAI-1) (Eriksson and Saldeen 1995). This, in turn, leads to weaker fibrinogen-polymerisation, thus making the clot more prone to lysis (Carlin et al. 1979).

Our group recently compared the effect of resuscitation with either 6% dextran-70 or 5% albumin in equipotent doses regarding plasma volume expansion on coagulation in a guinea pig haemorrhagic model. In that study, albumin reduced vWF levels to a larger extent than dextran, both immediately after resuscitation and 4 h later, but no difference in p-vWF GP1bA activity or in plasma fibrinogen levels could be detected (Schött et al. 2017). Dextran-70 resulted in a transient prolongation of thromboelastography (TEG) kinetics (K corresponding to ROTEM CFT) at the completion of resuscitation compared to albumin. The TEG reaction time (R, corresponding to ROTEM CT) and maximal amplitude (MA corresponding to ROTEM MCF) did not differ between the treatments at any of the time points (Schött et al. 2017). In vitro studies comparing dextran with other colloids indicated a deterioration in clot structure measured with another technique than ROTEM/TEG (Carr and Gabriel 1980; Carr 1986).

A recently published propensity-score matching study to evaluate the effect of dextran-70 on organ failure in severe sepsis did not show any signs of higher incidence of severe bleeding with dextran. Neither did it provide any evidence for detrimental effects of dextran-70 on 1979; Batlle et al. 1985). The mechanism behind this phenomenon is not fully understood and has even been observed with other artificial colloids. One theory is that macromolecules present in colloids bind to vWF, thus, accelerating its elimination (Treib et al. 1997; de Jonge et al. 1998). Additionally, dextran affects fibrinogen polymerisation by co-binding, increases the circulating levels of tissue type plasminogen activator (t-PA) and decreases the levels of fibrinolysis plasminogen activator inhibitor-1 (PAI-1) (Eriksson and Saldeen 1995). This, in turn, leads to weaker fibrinogen-polymerisation, thus making the clot more prone to lysis (Carlin et al. 1979).

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**Table 3 Multiplate values**

|                | Pre surgery | 500 colloid | 1000 colloids | Post surgery | Day 1 |
|----------------|-------------|-------------|---------------|--------------|-------|
| ADP (AUC)      |             |             |               |              |       |
| (Average)      | 89.1        | 76.6        | 95.8          | 81.9         | 77.2  |
| (Median)       | 86          | 87          | 96            | 84           | 75    |
| (Min-Max)      | (42–128)    | (44–110)    | (70–118)      | (44–108)     | (38–106) |
| COL (AUC)      |             |             |               |              |       |
| (Average)      | 81.7        | 77          | 99.2          | 85.1         | 82.2  |
| (Median)       | 84          | 62          | 95            | 82           | 74    |
| (Min-Max)      | (41–130)    | (40–118)    | (82–121)      | (38–123)     | (60–121) |
| TRAP (AUC)     |             |             |               |              |       |
| (Average)      | 118.7       | 125.4       | 138           | 128.4        | 107.8 |
| (Median)       | 117         | 124         | 131           | 130          | 108.5 |
| (Min-Max)      | (67–157)    | (78–170)    | (106–168)     | (87–171)     | (80–134) |
| Albumin Multiplate |       |             |               |              |       |
| Pre surgery    |             |             |               |              |       |
| (Average)      | 88.3        | 93          | 52.5          | 84.7         | 72.2  |
| (Median)       | 89          | 83.5        | 52.5          | 86           | 82    |
| (Min-Max)      | (63–116)    | (67–124)    | (22–83)       | (36–124)     | (32–99) |
| ADP (AUC)      |             |             |               |              |       |
| (Average)      | 83.7        | 97.7        | 63            | 86.9         | 77    |
| (Median)       | 78          | 82          | 63            | 88           | 85    |
| (Min-Max)      | (69–103)    | (60–149)    | (38–88)       | (43–149)     | (38–107) |
| COL (AUC)      |             |             |               |              |       |
| (Average)      | 132.4       | 124.7       | 84.5          | 123.6        | 102.2 |
| (Median)       | 137         | 120.5       | 84.5          | 126          | 88    |
| (Min-Max)      | (113–162)   | (85–176)    | (56–113)      | (76–176)     | (66–143) |

There were no significant intragroup changes (p > 0.05)

ADP adenosin phophate, COL collagen, TRAP thrombin receptor activating peptide for dextran and albumin at the different time points
kidney function, need for organ support or mortality (Bentzer et al. 2017).

Albumin is an endogenous protein with a molecular size of 39 kD. Albumin is the main component responsible for the colloid osmotic pressure in plasma and plays an important role in preserving intravascular volume (Statkevicius et al. 2016). Infusion with albumin derived from human plasma is frequently used for plasma volume expansion in the perioperative and intensive care settings. Albumin is an important transport protein and a free radical scavenger (Mortimer et al. 2014). Since albumin is derived from human plasma, it is relatively expensive and may facilitate transmission of infectious diseases. Albumin can also induce a dilutional coagulopathy and platelet defects (De Jonge and Levi 2001). Recent research has focussed on the glycocalyx, the vascular endothelium, and the role of plasma in maintaining its integrity. Albumin has, among other plasma proteins, been reported to have a protective effect on the glycocalyx (Schött et al. 2016). It has been speculated that the decrease in plasma concentration of markers of glycocalyx degradation after resuscitation with plasma as compared to crystalloids are largely secondary to differences in plasma volume and may not accurately reflect effects of FFP on the glycocalyx (Nelson et al. 2016).

We did not find any significant differences in the Multiplate AUC values for either of the groups. A similar find was detected with in vitro dilution with dextran 40 (Kam et al. 2017).

The Multiplate device measures platelet function using different activating factors corresponding to platelet receptors and has similar limitations as ROTEM. The Multiplate, therefore, cannot measure platelet dysfunction caused by vWF defects (Schött and Johansson 2013). Thus, in our case, we may suspect that the Multiplate failed to reflect a platelet dysfunction as dextran has a known effect on vWF. Studies have found that platelet counts of < 150 × 10⁹/l may influence the Multiplate, which should be considered in clinical settings (Stissing et al. 2011). The platelet count decreased more in the dextran group in the present study, but the Multiplate did not detect any differences between the albumin and dextran patients. In an earlier study on elective neurosurgery brain tumour resection from our centre, albumin decreased Multiplate aggregation more than HES (Li et al. 2015). An in vitro study with extreme colloid dilution (gelatine or HES; 60%) decreased Multiplate aggregation (Kind et al. 2013). Neither of these studies measured platelet counts. In fact, in the two patients receiving 1000 ml of albumin 5%, Multiplate aggregation was lower than in the six patients receiving 1000 ml of dextran-70 (Table 2). Whether this is a true effect of the colloid or is related to lower platelet counts needs to be studied with more patients. Currently, the European Society of Anaesthesiologists guidelines do not recommend perioperative Multiplate monitoring other than in patients with platelet inhibiting drugs (Kozek-Langenecker et al. 2014).

There were no statistically significant differences in bleeding or blood transfusions between the dextran or albumin groups, although the albumin group had less median bleeding (750 ml) as compared to the dextran

| Table 4 Standard laboratory (lab) values for Dextran and Albumin at the different time points |
|---------------------------------|-----------------|-----------------|-----------------|
|                                 | Pre surgery     | Post surgery    | Day 1           |
|                                 | n=7             | n=7             | n=6             |
| **Dextran routine lab**         |                 |                 |                 |
| Platelet count                  |                 |                 |                 |
| (Average)                       | 333.9           | 280*            | 218.8           |
| (Median)                        | 354             | 317             | 221             |
| (Min-Max)                       | (160–466)       | (114–432)       | (83–368)        |
| PT-INR                          |                 |                 |                 |
| (Average)                       | 0.99            | 1.3*            | 1.3*            |
| (Median)                        | 1               | 1.2             | 1.3             |
| (Min-Max)                       | (1.1–1.7)       | (1.1–1.5)       |                 |
| aPTT                            |                 |                 |                 |
| (Average)                       | 25.6            | 366*            | 31.6*           |
| (Median)                        | 25              | 31              | 30              |
| (Min-Max)                       | (24–28)         | (27–61)         | (28–42)         |
| Fibrinogen                      |                 |                 |                 |
| (Average)                       | 4.6             | 2.0*            | 3.3             |
| (Median)                        | 4.2             | 1.9             | 3.35            |
| (Min-Max)                       | (3.2–8.2)       | (1–3)           | (1.6–5.1)       |
| **Albumin routine lab**         |                 |                 |                 |
| Platelet count                  |                 |                 |                 |
| (Average)                       | 328.4           | 267.7           | 228.4*          |
| (Median)                        | 277             | 241             | 188             |
| (Min-Max)                       | (238–483)       | (143–422)       | (124–381)       |
| PT-INR                          |                 |                 |                 |
| (Average)                       | 1               | 1.3             | 1.3*            |
| (Median)                        | 1               | 1.3             | 1.4             |
| (Min-Max)                       | (1–1)           | (1–1.5)         | (1.1–1.5)       |
| aPTT                            |                 |                 |                 |
| (Average)                       | 27.1            | 30.7            | 32.3            |
| (Median)                        | 27              | 29              | 33              |
| (Min-Max)                       | (25–29)         | (25–41)         | (26–37)         |
| Fibrinogen                      |                 |                 |                 |
| (Average)                       | 4               | 2.1*            | 3.3             |
| (Median)                        | 4               | 1.8             | 3.2             |
| (Min-Max)                       | (2.9–5.4)       | (1.4–4.2)       | (2.5–4.2)       |

*Significant (p < 0.05) intragroup change from the respective pre surgery value
group (900 ml) and needed fewer transfusions of PRBC. The low number of patients receiving colloids in the present study (n = 14 in total) and the variation in pre-operative coagulation profiles—especially preoperative ROTEM indicating hyper-, normo- and hypocoagulation (Table 2, Fig. 1)—blur the interpretation of whether it is the colloid-induced coagulopathy, the difficulties in resection of tumours or the difficulties in obtaining surgical haemostasis that increases bleeding and the higher transfusion rate in the dextran group. The large ranges in bleeding (200–3500 ml in the dextran group and 200–1100 ml in the albumin group) are also a confounding factor impeding the interpretation of our results. A previous study from our centre has implemented the nomenclature *dilutional capacity* to refer to the above-described preoperative coagulation strength measured with ROTEM and specific coagulation factor analyses (fibrinogen, FXIII activity, thrombin-antithrombin complex [TAT] and plasmin-α2-antiplasmin complex [PAP]) to individualise the colloid use to avoid colloid induced dilutional coagulopathies (Nilsson et al. 2016). This approach should be tested, but it requires at least a preoperative ROTEM/TEG testing, currently not recommended by the European Society of Anaesthesiologists (ESA) guidelines for perioperative bleeding management (Kozek-Langenecker et al. 2014).

Perioperative TEG hypercoagulation has also been suggested to reflect the degree of malignancy of ovarian masses (Amirkhosravi et al. 2013). Dextran has been suggested to reduce thromboembolism in cancer surgery (Foster et al. 2017). Dextrans have also been used in vascular surgery to diminish the risk of stroke associated with carotid endarterectomy and in “difficult” infrageni- cular lower extremity bypasses (Abir et al. 2004). However, a recent retrospective study found no reduction in perioperative stroke, but more cardiac failure and myocardial infarction after dextran (Farber et al. 2013).

**Limitations**

There were several limitations in this study. It was an open observational screening study of coagulation changes with two fluid regimes, and it was non-randomised and not blinded. It was a pilot study of a problematic group of patients undergoing major gynaecological cancer surgery that had a risk for major bleeding and often showed rapid fluid shifts intra- and postoperatively that demanded a high level of postoperative surveillance. It was underpowered to detect a correlation between ROTEM, laboratory and Multiplate analyses and clinical parameters such as bleeding, the need for transfusions and postoperative complications. The ROTEM or Multiplate system results might not always correlate with clinical bleeding, as there were no blood flow/shear stress, effects of endothelial activation/inhibition or effects of the glycocalyx layer that interacted with coagulation and platelet activation in vivo. The study also had a small sample size, reflecting the limited period of patient recruitment time, but our aim is to proceed with this study up to 60 patients.

**Conclusion**

Standard plasma-based coagulation tests, platelet count and whole blood viscoelastic clot structure are affected by 6% dextran-70 to a greater extent than by 5% albumin, but platelet aggregation is not. Future studies should use more advanced haemodynamic monitoring to assess isovolemic plasma volume expansion with dextran and whether this affects haemostasis to a lesser degree.

**Abbreviations**

AUC: Area under the curve; 6S: Scandinavian Starch for Severe Sepsis/Septic Shock trial; A10: Clot formation at 10 min; AA: Alpha-angle; ADP: Adenosin phosphate; APPT: Activated partial thromboplastin time; BW: Body weight; CFT: Clot formation time; COL: Collagen; CT: Clotting time; F: Factor; FASS: Pharmaceutical Specialities in Sweden; HES: Hydroxyethyl starch; INR: International normalised ratio; K: Kinetics; kD: kiloDalton; MA: Maximal amplitude; MAP: Mean arterial pressure; MCF: Maximum clot formation; PAI-1: Plasminogen activator inhibitor-1; PAP: Plasmin-α2-antiplasmin complex; POC: Point of care; PRBC: Packed red blood cells; PT: Prothrombin time; R: Reaction time; ROTEM: Thromboelastometry; TAT: Thrombin-antithrombin complex; TEG: Thromboelastography; t-PA: Tissue type plasminogen activator; TRAP: Thrombin receptor activating peptide; vWF: von Willebrand factor

**Funding**

Lund University funding for Ulf Schott (ISEX-ALF).

**Availability of data and materials**

Peter Bansch/Ulf Schott can provide original data in excel files, graph prints from ROTEM/Multiplate devices.

**Authors’ contributions**

PB and US planned the study, wrote and processed the ethical application and tutored DH (medical graduate) and JS (MD, specialist in training). All authors contributed to sampling and ran analyses on point of care devices. DH processed the sampling and its transport to the routine coagulation analyses at the Department of Clinical Chemistry. All authors performed analyses, statistical evaluation with tables and figures and wrote the draft. All authors read and approved the final manuscript.

**Ethics approval and consent to participate**

Yes, see “Methods” section.

**Consent for publication**

Yes, see “Methods” section.

**Competing interests**

The authors declare that they have no competing interest.

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Received: 17 April 2018 Accepted: 30 July 2018 Published online: 06 September 2018

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