Thrombotic Occlusion of a Microvascular Anastomosis in a Resistance to Activated Protein C (APC) Patient with Incomplete Wound Healing after High Doses of Ascorbic Acid (Vitamin C)

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ABSTRACT

A 45-year-old woman underwent a delayed breast reconstruction with a free Deep Inferior Epigastric Perforator Flap (DIEP flap) with total flap failure on the fourth postoperative day. Hematological investigation to exclude thrombophilia revealed a resistance to activated protein C (APC) with a factor V Leiden heterozygous mutation. The postoperative course was further complicated by delayed wound healing probably due to ascorbic acid (Vitamin C) related cytotoxic activity to fibroblasts. The surgeon must be aware of the use of preoperative nutritional supplement administration among patients. Future cost-effectiveness analyses should be made to warrant preoperative thrombophilia screening to prevent free flap failures.

Keywords: Thrombotic Occlusion; Vitamin C; Activated Protein C

1. Case Report

A 45-year-old woman underwent a delayed breast reconstruction with a free Deep Inferior Epigastric Perforator Flap (DIEP flap) in our department. The surgical history reported an appendectomy 10 years ago and a mastectomy 2 years ago for an invasive ductal carcinoma with a positive sentinel node biopsy followed by axillary lymph node dissection. No postoperative complications occurred during these procedures.

She reported a superficial vein thrombophlebitis in one leg during her chemotherapy.

Her medicine history reported no medication administration during the last 6 months preoperatively and no smoking history.

During the surgical procedure, we found an arterial spasm after the microsurgical anastomosis. Exploration of the arterial anastomosis revealed no thrombus and a new second anastomosis was performed without further surgical complications.

Postoperative intravenous Piracetam (Nootropil; 12 gram/24 hours) and subcutaneous Nadroparine injections (Fraxiparine; 2850 IE anti-Xa (= 0.3 ml/24 hours) were administered.

Postoperative flap monitoring during the first three postoperative days revealed no signs of flap ischemia or venous congestion.

On the fourth postoperative day however, routine clinical evaluation of flap vitals showed severe congestion of the whole flap (Figure 1). Emergency surgery with re-exploration revealed a venous thrombus with severe congestion injury of the flap tissues. Thrombectomy was performed after which the flap was infused with a thrombolytic agent (Urokinase). The vascular microanastomosis was revised and an arterial interposition graft was used to support the venous anastomosis.

A clinical re-occlusion of the vascular anastomosis was found 6 hours after the revision.

We decided to amputate the flap in consultation with the patient.

Partial central abdominal wound dehiscence started on day 5 post-operatively with abnormal amounts of wound fluid drainage based on fat necrosis (Figure 2). Secondary infection with non-fragilis bacteriodes was found and intravenous amoxicillin with clavulanic acid was started. Progressive necrosis and dehiscence happened until 14 days after the operation after which wound stabilization occurred.

Hematological investigation after the flap amputation, to exclude thrombophilia, revealed a resistance to activated protein C (APC) with a factor V Leiden heterozygous
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2. Discussion

2.1. Resistance to Activated Protein

Resistance to activated protein C (APC) is the most common genetic risk factor associated with venous thrombosis. Most cases are due to a point mutation in the factor V gene (factor V Leiden, FVL), which subsequently prevents the cleavage and disruption of activated factor V by APC and thus promotes ongoing clot development. In approximately 3% - 8% of white adults, this mutation is heterozygous, conferring a 5-fold increased lifetime risk of venous thrombosis compared with the general population [1].

Irreversible thromboses after microvascular anastomosis due to hereditary thrombophilia have been described in the literature [2,3]. Furthermore, we found a former comparable free DIEP flap failure after an irreversible venous thrombosis in another patient with an APC resistance operated in our department. Future cost-effectiveness analyses should be made to warrant preoperative thrombophilia screening to prevent free flap failures.

2.2. Ascorbic Acid (Vitamin C)

The former appendectomy and mastectomy wounds in this patient healed without complications and therefore we link the abdominal wound dehiscence to her 4 months of highly dosed Vitamin C administration.

Pharmaceutical dosis of ascorbic acid have been reported to exert cytotoxic and anticancer activity in vitro and in vivo [4]. Increased ascorbic acid concentrations are found in the extracellular fluid in animal models after intravenous administration of pharmacologic concentrations of vitamin C while substantial lower concentrations are found in blood [5]. One proposed working mechanism involves direct cytotoxicity mediated by accumulation of ascorbic acid radicals and hydrogen peroxide in the extracellular environment of tumor cells but also to fibroblasts which may contribute to incomplete wound healing [6,7]. This could be an explanation for the delayed wound healing in our patient.

Secondly, high doses of ascorbic acid suppress NO (nitric oxide) generation. The endothelium of blood vessels uses nitric oxide to signal the surrounding smooth muscle to relax, thus resulting in vasodilatation and increasing blood flow. This angiostatic effect of vitamin C...
may explain the arterial spasm after the microsurgical anastomosis during the surgical procedure [4]. However, the hypothesized tissue accumulation of vitamin C in this patient needs to be further proven by laboratory testing in human models. Nevertheless, we suggest stopping preoperative administration of high doses of nutritional supplements.

3. Conclusion

The surgeon must be aware of the use of preoperative nutritional supplement administration among patients. Future cost-effectiveness analyses should be made to warrant preoperative thrombophilia screening to prevent free flap failures.

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