Neutrophil Extracellular Trapping Role in Cancer, Metastases, and Cancer-Related Thrombosis: a Narrative Review of the Current Evidence Base

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Abstract

Purpose of Review Neutrophil extracellular trap (NET) formation is a newly discovered, reactive oxygen species-dependent regulated process, whereby neutrophils degranulate and extrude genetic material, after engulfing various infectious or neoplastic antigens, culminating in a measurable serologic footprint. Recent research has highlighted the involvement of NETs in cancer and cancer-related pathologies. We review the role of NET formation in cancer biology, prognosis and potential therapeutic modulators.

Recent Findings Elevated NET levels are associated with cancer metastasis and may be modified by some anaesthetic-analgesic techniques during tumour resection surgery. It promotes tumour cell migration, angiogenesis and hypercoagulability. Although there are potential anti-NET formation therapeutics available, their role has not been formally assessed in cancer patients.

Summary Limited available evidence suggests an association between elevated NET expression and cancer metastasis, but its validity as a prognostic indicator for cancer-related outcomes is inconclusive. Further observational and interventional studies are warranted to comprehend the potential prognostic and therapeutic role of NETs in cancer.

Keywords Neutrophil extracellular trapping · NETosis · NETs · White blood cells · Neutrophils · Cancer · Metastases · Thrombosis

Introduction

Neutrophil extracellular trap (NET) formation is a reactive oxygen species (ROS)-dependent regulated process, whereby neutrophils degranulate, releasing intracellular and nuclear contents which form an extracellular snare with a role in bacterial or neoplastic cell trapping and killing. This process bequeaths a serologic footprint which may be detected many years later. The term NETosis has been previously used for describing this phenomenon. Recently, NETosis was defined as being a ROS-dependent regulated cell death, specific to cells of haematopoietic origin and associated with NET release [1]. It is recommended to avoid using the term NETosis in those instances where there is no evidence of cell death associated with NET formation. The term NET extrusion may also be used for NET formation without cell death. These two terms will be used interchangeably for the remaining of this article.

NETs are rich in cell nuclear components (e.g. DNA, histone proteins). They also contain granular and cytoplasmatic proteins or enzymes (myeloperoxidase (MPO), neutrophil elastase (NE)) [2] and antimicrobial and proinflammatory proteins [3]. A defining characteristic of NET generation is the translocation of genetic material from the nucleus to the extracellular environment [4]. Although NET formation was initially described over 20 years ago, it has only emerged as a current topic of interest since 2014 [5, 6]. Since then, it has been identified as a factor...
in many physiological and pathological processes like immunity, sepsis, inflammation, cancer metastasis, cancer-related thrombosis, and more recently in severe acute respiratory syndrome coronavirus 2 (SARS-CoV 2) infection [7–11].

In this review, we summarise the current understanding of the multifactorial role of NETs in tumour cell biology, metastatic development and venous thromboembolism (VTE) occurrence. We also review pharmacological agents, including anaesthesia technique, with a potential role in NET modulation.

**Mechanisms of NET Formations**

Classically, neutrophils are recognised as the mainstay of the innate immune system, delivering chemotaxis, endothelial adherence, phagocytosis, oxidative burst and toxic granule dependent microbial killing [4]. Important steps of NET formation include chromatin decondensation, nuclear membrane rupture, nucleoplasm spillage into the cytoplasm, cell membrane destruction and cell death have been described. This has been described both in vitro and in vivo.

Suicidal NET formation is NADPH oxidase (NOX)–dependent and involves chromatin decondensation, nuclear envelope disintegration, plasma membrane perforation, cell lysis and tumour cell or bacteria fixation to the DNA structures [12, 13]. Subsequent work revealed the role of ROS, NE and MPO in NET release [2]. The ROS triggers the release of NE and MPO from the cytoplasmic granules and promotes the MPO-mediated proteolytic activity of NE. The NE and MPO are translocated to the nucleus, which facilitates histone degradation chromatin decondensation [2].

This contrasts with “vital NETosis”, a more rapid process in which neutrophils became anuclear cells still capable of survival and of engulfing antigens [4, 14]. Currently, three types of vital NET formation have been described (Table 1).

Suicidal and vital NET formation terminology is being gradually replaced by NOX-dependent and NOX-independent NET production. NOX-dependent or NOX-independent pathway is activated in response to specific triggers. Lipopolysaccharide (LPS) and bacteria stimulate the NOX-dependent pathway leading to nucleus breakdown [2] and activation of a specific set of kinases (ERK1, ERK2, Akt, P38, Src) [17**].

Bacterial or neoplastic cell products and uric acid crystals activate the NOX-independent pathway, leading to the formation of mitochondrial ROS [18]. This acts on several kinases. The activation of each pathway’s kinases leads to transcriptional firing, which enables chromatin decondensation [6]. In the NOX-independent pathway, peptidyl arginine deaminase type 4 (PAD4)-mediate the production of chromatin networks by the cancer cells themselves [19, 20]. There is also evidence that NET formation can take place in a PAD4-independent fashion [21] particularly in the NOX-dependent pathway [18, 22].

**Measuring NET Expression**

The production of NETs creates a footprint, by which NET formation can be detected. The various components of NETs (e.g. histones, DNA, enzymes) and the molecules involved in NET expression (e.g. PAD-4) serve as potential targets for measuring this process. NET expression can be quantified directly by microscopy [23], flow-cytometry or real-time imaging or indirectly, using the enzyme-linked immunosorbent assay (ELISA) technique mainly for serological measurements [24]. MPO, NE, and citrullinated histone 3 (citH3), among others, represent measurable components or markers of NETs [25, 26, 27*], but may not be specific. For example, NE and MPO can be released during degranulation without NET formation [28]. It has been proposed that measuring conjugates like MPO/circulating free DNA (cfDNA) and citH3 could be more specific for NETs than cfDNA alone [29].

**NETs and Cancer**

Elevated NET levels are associated with the diagnosis of cancer [4, 10, 17**, 27*]. NETs contribute to primary tumour growth [9], metastasis [30] and cancer-related thrombosis [31] through various mechanisms (Fig. 1). NETs associated enzymes are involved in cancer growth by stimulating phosphatidylinositol-3 kinases (PI3Ks) (e.g. NE) and matrix destruction (e.g. matrix metallopeptidase 9 (MMP-9)) [36, 37]. In breast cancer cells in vitro, NET expression upregulates gene expression for several factors related to

**Table 1** “Vital” NET formation subtypes

| Subtype | Manuscript | Triggers |
|---------|------------|----------|
| NETs containing mitochondrial DNA | Yousefi et al. [15] | Granulocyte/macrophage colony-stimulating factor (G-CSF), TLR4 or complement factor 5a (C5a) receptor stimulation via a ROS mechanism |
| NETs (NOX-independent) | Pilszczek et al. [16] | *Staphylococcus aureus* |
| NETs (NOX-independent) | Yipp et al. [14] | Lipopolysaccharide via TLR-4 receptor on platelets. Requires platelets/neutrophils crosstalk |
pro-metastatic properties of breast cancer [38]. Also, cancer cell biology and NET formation share common elements of ROS signalling pathways [39] and reliance on similar kinases (e.g. p38) [17, 40]. For example, stimulation of toll-like receptor 4 (TLR4) receptor (e.g. by LPS) induces NET formation [32] with a role in host immunity. High mobility group box protein 1 (HMGB1) protein is involved in many steps of the protumorigenic pathway [41] and is produced by neutrophils on degranulation/NET formation. This protein seems to be one of the NETs associated chemotactic factors responsible for the NET-induced tumour cell invasion [26]. Other proteins released during NET production like MMP-9 and NE have been shown to be involved in tumour progression, creating an adequate environment for tumour cell seeding and growth [26]. Despite some evidence that in vitro cancer-induced NET formation is a suicidal process [42], it remains unclear if the in vivo process is vital or suicidal [32].

In an in vitro model of surgery, it was shown that NETs stimulated Kupffer cells to produce proinflammatory cytokines directly associated with cancer progression (interleukin 6 (IL-6), tumour necrosis factor-alpha (TNF-alpha) and chemokine CXC motif ligand 10 (CXCL-10)) [26, 43]. This was alleviated by DNase and PAD4 inhibition [26]. Although the full list of stimuli of NET production in cancer is unknown, there is evidence for the circulation of pro-NET formation factors. Neutrophils from healthy donors were stimulated to produce NETs when incubated with cancer patient’s plasma [44].

**Tumour Cell-Neutrophil Interplay in NET Formation**

Tumour cell-neutrophil-platelets-vascular endothelium interactions are essential for cancer cell survival and the development of conditions which facilitate metastasis. These
processes have been recently summarised [32••]. Neutrophils are subject to modulation form both circulating tumour cells and factors released by the solid tissue tumours through extracellular vesicles (EVs). Some of the identified modulators produced by the primary tumours are tissue factor (TF), granulocyte colony-stimulating factor (G-CSF), HMGB1, IL-8 and ROS. The interaction between tumour factors, activated platelets and endothelium leads to neutrophil activation, NET formation and adhesion of the neutrophils to the endothelium [45]. In breast cancer, increased G-CSF is associated with increased metastases [46]. In a murine model of breast cancer and insulinoma, it was shown that cancer-associated heart, kidney and vascular dysfunction, which caused increased NETs, was improved by neutrophil depletion, G-CSF inhibition and DNAse administration [47]. In this model, vascular NET development was mediated by increased levels of intercellular adhesion molecule 1 (ICAM-1), vascular cell adhesion protein 1 (VCAM-1), IL-1Beta, IL-6 and chemokine CXCL1. These observations may explain the distant organ dysfunction seen in cancer.

Extracellular vesicles are membrane-enclosed structures produced by cells (e.g. during cell stress response) and could contain specific cytoplasmic solid and soluble components specific to the parent cell [32••]. Tumour EVs can directly trigger NET formation after they have been phagocytosed by neutrophils. They have been found to be present in patients with breast cancer and to predict therapeutic failure and poor prognosis [48]. Some of the constituents of EVs (e.g. interleukins and G-CSF) are NET formation mediators [49]. In a murine model (BALB/c mice) injected with 4T1 breast cancer cells, the EVs of 4T1 cells triggered NET formation and when administered intravenously contributed to accelerated venous thrombosis [50]. In a mouse model of lung tumours postintravenous B16 melanoma cell injection, the authors showed that the tumour cell EVs were ingested by neutrophils which could have pro-NET generation effects.

In addition, NETs act like “fishing nets” capturing circulating tumour cells which generate further endothelial damage facilitating metastasis [17••]. Key ligand/neutrophils receptor interactions (e.g. HMGB1-TLR4/receptor for advanced glycation end products (RAGE)) are essential in this scenario [32••] and could develop into therapeutic targets. Although the IL-8/CXC chemokine receptor 1/2 (CXCR1/2) interaction is beneficial in acute infection [52], it seems to be detrimental in cancer. In a murine model of lung cancer and concomitant sepsis, intravascularly deposited NETs captured lung cancer cells and were associated with gross metastatic burden at 2 weeks [53]. The metastatic burden was attenuated by NET formation inhibitors (e.g. DNAse, neutrophil elastase inhibitor (NEi)). In a murine model of murine colon cancer (CT-26) peritoneal metastasis, decreasing the circulating neutrophils or disrupting NET formation by DNAse markedly decreased the number of metastases [30]. In the same study, mice and humans were shown to have significant NETs in peritoneal metastases from colon cancer, and blocking the integrins abolished NET-induced cancer cell adhesion.

Cancer cell NET formation occurs by two pathways: pro-tumorigenic effects of NETs and cancer cells’ NET-inducing capacity. In a study involving patients with pancreatobiliary malignancy and the pancreatic cancer cell line AsPC-1, cells themselves and their conditioned medium were strong promoters of NET formation. In this study, ROS inhibition did not block NET formation but prostaglandin E1 (PGE1) did, suggesting a role of cyclical adenosine monophosphate (AMP) in NET formation [54]. It was shown that HMGB1 protein is released from NETs and serves as a ligand on the TLR4 receptor on the surface of cancer cells [26, 27•, 55].

The age of the neutrophil is relevant for both NET formation and cancer biology [56]. Interestingly, the more immature neutrophils have a reduced NET formation activity via reduced ROS production and increased mitotic activity [57].

**NETs and Cancer-Associated Thrombosis**

Endothelial cells and platelets require prior activation before triggering NET formation [58]. Platelets can be activated by the histones in NET structure [59] and can directly stimulate NET formation, interacting with the neutrophil via the selectin P (P-SEL)/selectin P receptor 1 (PSGL-1) [32••]. NETs are rich in citrullinated histone 3 (citH3), NE, and MPO, all triggering further platelet activation and contributing to the thromboembolic phenomenon associated with cancer. In addition to the previously known mechanisms of platelet activation, an in vitro study using AsPC-1 pancreatic cell line suggested platelet activation may be driven by the tumour cells directly via TF expression [60], which could potentially be a constituent of the tumour EVs [32••]. HMGB1 factor and TLR4 receptor are used by the activated human platelets to trigger NET formation [58]. Furthermore, DNA activates the intrinsic coagulation pathway [61]. Jung et al. proposed that the DNA contained in NETs could be the trigger for NET-associated thrombin generation and subsequent cancer-related thrombotic phenomena [27•].

NETs have been identified in patients with thrombotic phenomena (i.e. stroke). In a prospective, observational case-control study, patients were divided into two groups based on the serum troponin levels [62]. High-sensitivity troponin (hsTnT) was associated with a higher risk of malignancy. Autopsies in these patients showed widespread microthrombosis containing citH3, a biomarker for NETs. A procoagulant state with increased NET serum levels was seen in patients with elevated hsTnT compared to that in patients with normal levels (p < 0.001). CitH3 was also correlated with thrombin-antithrombin complex and soluble P-SEL providing a further link between NETs and the prothrombotic state in cancer patients.
A similar prothrombotic state was found in a study of patients with pancreaticobiliary malignancy, who had a high level of circulating NET markers (histone-DNA complexes and cfDNA) and hypercoagulability markers \[27^\text{•}\]. Also, hypercoagulability markers were higher in patients with more advanced cancer (stage IV vs stage I/II). In the same study, using the AsPC-1 pancreatic cancer cell line, the investigators also documented that antithrombin significantly blocks NET formation.

**NETs as a Prognostic Marker**

Plasma taken from treatment-naive patients with lung or upper GI adenocarcinoma and compared to the plasma of healthy individuals showed an elevation in NET expression (measured as MPO) in the cancer patient cohort compared to that of the control group (p = 0.03). This study also showed significant elevation in NET levels in patients with advanced cancer (stages III & IV oesophagogastric, stages II & III lung) compared to patients with a less advanced diagnosis (stages I & II oesophagogastric and stage I lung) (p = 0.01) \[63^\text{•}\]. NET expression was also higher in patients with early-stage head and neck cancer compared to that in healthy control participants of the same age \[64^\text{•}\]. As part of the same study, it was shown that neutrophils of these cancer patients released more NETs compared to those of the healthy control cohort. The increased NET expression in cancer patients and the increased release of NETs from neutrophils in vitro support the hypothesis that NETs may be a biomarker for cancer.

Although most evidence points towards a deleterious effect of NETs in cancer, a multi-omics study on ovarian cancer patients found that NET formation correlated with better outcome \[25\]. Increased CRP is typically associated with poor prognosis in this type of cancer, whilst the S100A8 protein is released by neutrophils upon local NET generation. The S100A8/CRP abundance ratio significantly stratified patients and the increase in the ratio correlated with favourable survival.

The opposite was observed in patients with colorectal cancer, with an in vitro increase in NET expression in response to stimulation of systemic neutrophils. This was associated with more postoperative complications (Clavien-Dindo classification ≥3) and prolonged hospital stay (>5 days). \[23\].

NET formation has been predictive for decreased disease-free survival. In a cohort of 50 colorectal patients undergoing curative liver metastases resection, it was observed that increased postoperative NET formation was associated with over fourfold reduction in disease-free survival \[26\]. Similar findings occurred in a murine model (C57BL/6J WT mice or PAD4−/− mice) of surgical stress employing liver ischemia-reperfusion, where NET expression was associated with an accelerated development of metastatic disease (MC38 or MC38/Luc cancer cells) \[26\]. These effects were mitigated by inhibiting NET formation with DNase (68% reduction in hepatic tumour burden) or PAD4 inhibition.

In a study of 60 patients with various adenocarcinomas, squamous cell carcinomas, melanomas, and glioblastomas, there was a threefold increase in the median plasma citH3 (NET biomarker) levels in patients with advanced cancer compared with age-matched healthy individuals or severely ill patients without cancer \[65\]. Furthermore, plasma levels of IL-8, IL-6, TNF-alpha, IL-1beta and G-CSF (all involved in NET production) were all elevated in cancer patients. Levels of citH3 above the 75th percentile were associated with a twofold short increased short-term mortality. Besides, these patients had a significantly higher number of peripheral neutrophils, neutrophils positive for intracellular citH3, plasma MPO, and MPO-DNA complexes, suggestive of neutrophil activation and NET extrusion. In a systematic review of 30 studies where the authors examined the relationship between perioperative inflammation, NET expression and metastasis in colorectal cancer, the presence of a preoperative systemic inflammatory state but not sepsis was associated with increased cancer recurrence \[66\].

Cancer cells can hijack neutrophil function, inducing NET formation independently of direct tumour cell-neutrophil interaction (via G-CSF priming) and could be a marker of tumour aggression. In an in vitro study, NETs were found in 16/20 of the primary breast tumours and in 13/19 metastatic lung lesions. Lung metastases of triple-negative breast cancer had the highest level of NET expression when compared to luminal breast cancer \[42\]. Interestingly, these in vitro experiments revealed a lytic (i.e. suicidal) NET formation pathway and concluded that NETs do not promote tumour cell extravasation, but rather mediate the expansion of already disseminated cells.

**NETs Promote Angiogenesis**

NETs are promoters of inflammation via NE and MPO, which could damage the endothelium, triggering further IL-8, ROS and platelet activation, in turn further stimulating neutrophil recruitment and NET release \[32^\text{•}\]. In addition, in vitro, histones are stimulators of angiogenesis when incubated with endothelial cells \[27^\text{•}\]. As they are a major component of NETs, it is reasonable to conclude that NETs can promote angiogenesis too. The entire effect was blocked by histone chelating agents (e.g., heparin, polysialic acid (PSA)).

**Does Anaesthesia Technique and the Perioperative Surgical Stress Impact on NET Formation?**

The effects of surgical stress and anaesthetic technique on NET expression have been assessed in vivo and clinically, with intriguing results.
| Substance | Manuscript | Experimental model | Findings | Comments |
|-----------|------------|--------------------|----------|----------|
| Anthracyclines (e.g. epirubicin, daunorubicin, doxorubicin and idarubicin) | Khan et al. [74] | Human neutrophils, in vitro | Both NOX-dependent and NOX-independent NET expression were blocked. | Anthracyclines’ effect may be mediated by alteration of transcription initiation. |
| Anti-GCSF antibodies | Park et al. [42, 75] | Murine model. 4T1 breast cancer cells | Reduced cancer cell–induced NET expression and metastases. | In humans, these could be difficult as GCSF is commonly used as a treatment of neutropenic sepsis in cancer patients [42]. |
| Anti-HMGB1 antibodies | Tohme et al. [26] | In vitro. MC 38 colorectal cancer cells | NET-mediated tumour cell proliferation, invasion and migration were significantly decreased (p < 0.001). | Leads to MAP kinase pathway inhibition. |
| Antithrombin III | Ishikawa et al. [76] | In vitro. MC 38 colorectal cancer cells | Reduced mortality, lung neutrophil infiltration, expression of NETs, HMGB1 protein and CXCL1 and 2. | In addition to the anticoagulation effects, antithrombin III has also anti-inflammatory effects. |
| COX inhibitors | Tilgner et al. [69] | In vivo, LPS-induced acute lung injury model in C57Bl/6 mice | Aspirin decreased neutrophil infiltration and NET formation. | Although a drug used as an anti-parasitic, this could be useful in NET expression modulation in diabetic wound healing and cancer potentially. |
| Diethylcarbamazine | Segoviano-Ramirez et al. [77] | In vitro. polymorphonuclear cells (PMNs) from healthy and type 2 diabetes mellitus patients | NET formation was significantly inhibited both in healthy (p ≤ 0.000001) and DM2 (p ≤ 0.000477) volunteers. | Among all other modulators, DNase has the largest use in various experimental models and is already in clinical use for CF treatment. It may likely be among the first drugs to be used in cancer for outcome control. |
| DNase | Jung et al. [27*] | In vitro. AsPC-1 pancreatic cancer cells | Significant inhibition of AsPC-1-induced NET formation. | Aspirin decreased neutrophil infiltration and NET formation via CCL5 chemokine and PAD4 inhibition. |
| Diethylcarbamazine | Jung et al. [27*] | In vitro, murine model. MC 38 colorectal cancer cells | Significant inhibition of tumour adhesion, the development and growth of metastatic disease. | Among all other modulators, DNase has the largest use in various experimental models and is already in clinical use for CF treatment. It may likely be among the first drugs to be used in cancer for outcome control. |
| Park et al. [42] | Murine model. 4T1 breast cancer cells | DNase-coated nanoparticles decreased lung metastatic burden. | Preliminary in vitro and animal results suggest DNase reduces NET formation, with effects on tumour cell adhesion, tumour burden and metastasis [26, 42]. |
| Kolaczkowska et al. [78] | In vitro. human neutrophils | Significant inhibition of AsPC-1-induced NET formation. | | |
| Gasdermin D inhibitor LDC7559 | Sollberger et al. [79] | In vitro, human neutrophils | Gasdermin D is required for NET expression and acts upstream of NE. LDC7559 reduced histone H3 protein processing. | Gasdermin D is a pore-forming protein involved in the release of granular proteins required for NET expression and facilitates NE translocation to the nucleus. Also forms membrane pores releasing the NETs in the extracellular space. |
| Heparin and polysialic acid (PSA) | Jung et al. [27*] | In vitro, AsPC-1 pancreatic cancer cells | Blocked histone-induced angiogenesis. | These two substances have in common a highly negative electric charge which binds to positively charged histones, leading to blockade of NET-induced cancer cell migration and angiogenesis [27, 80]. |
| HMG-CoA reductase inhibitors | Sapey et al. [81] | In vitro. neutrophils from patients receiving high-dose simvastatin for community-acquired pneumonia | Significant reduction in NET expression and NE activity on day 4. No difference in mortality and length of stay | The result of these studies looking at the NETs role in response to bacteria is very different. Keeping in mind the different experimental models, translation of anti-NET therapies from bench to bedside may prove difficult and may not be |
| Chow et al. [82] | In vitro. Human and mice PMNs, cultured with various statins: mevastatin, simvastatin, lovastatin and | Statins boosted the production of anti-bacterial DNA-based extracellular | | |
Table 2 (continued)

| Substance | Manuscript | Experimental model | Findings | Comments |
|-----------|------------|--------------------|----------|----------|
| Lidocaine | Galos et al. [67] | Patients undergoing primary breast cancer resection | fluvastatin. The NET formation was quantified in response to bacterial exposure. Lidocaine infusion decreased expression of citH3 and MMP-3. | associated with a clinical effect. This could be the case in cancer too. |
| NADPH oxidase 2 (NOX2) inhibitors (e.g. apocyanin) | Park et al. [42] | In vitro. Murine model. 4T1 breast cancer cells | NOX2 inhibition reduced NET formation and neutrophil-promoted cancer cell invasion. | This study offers the premises for a large trial regarding the impact of lidocaine on the metastasis. NADPH oxidase is a critical enzyme as shown already in the NET production process. Although inhibitors against it are available and represent a potential tool for blocking NET production, it is not feasible as this enzyme is involved in critical processes like bacterial killing, phagocytosis and degranulation [42]. |
| PAD4 inhibition (e.g. YW4-03, Cl-amidine) | Tohme et al. [26] | In vitro and a murine model. MC 38 colorectal cancer cells. PAD4-deficient mice | In vitro, PAD4 inhibitors or the use of PAD4-deficient neutrophils block the adhesion of tumour cells to a monolayer of NETs expressing neutrophils. Significantly reduced the tumour cell growth, p < 0.0001 Levels of cit-H3 in the tumours after 3 weeks were absent in PAD4−/− mice and mice receiving YW4-03. | PAD-4 inhibition has been associated with preventing NET formation in animal models, but its effect on metastasis will be difficult to predict due to the short half-lives of the current commercially available inhibitors [42]. |
| Park et al. [42] | Murine model. 4T1 breast cancer cells | It decreased NET formation (p = 0.02). No effect on tumour cell migration | | |
| PGE1 | Jung et al. [27] | In vitro, AsPC-1 pancreatic cancer cells | Inhibited cancer cell–induced NET formation, by decreasing the intracellular calcium content | PGE 1 has been used clinically for neonatal duct–dependent disease, sexual dysfunction, critical limb ischemia due to its anti-inflammatory, vasodilative and anti-platelet effects [83]. Both extra and intracellular alkaline Ph have been linked with cancer promotion [85]. Also, Ph-sensitive channels could become a therapeutic target in some cancers [86]. The alkaline pH modulates the NET formation as enzymes like PAD4 and NE prefer an alkaline environment and processes like mitochondrial ROS and PAD4-mediated citrullination are augmented by alkaline pH with an impact on both NOX-dependent and NOX-independent processes [84, 87]. |
| Ph | De Souza et al. [84] | In vitro. Human neutrophils | Raising the pH promotes intracellular calcium influx which leads to increases PAD4 activity, citrullination of histone, histone cleavage and NET formation. | |
In an observational cohort study evaluating the metastatic potential of surgical stress-induced NET formation, liver ischemia-reperfusion (I/R) injury increased NET expression. Serum was obtained day 1 postoperatively for patients who underwent hepectomy for metastatic colorectal cancer. NET expression (measured by serum MPO-DNA complexes) was significantly increased in patients who underwent major liver resection (3 or more segments removed) with unavoidable liver I/R injury when compared to patients who underwent smaller resection or to healthy controls. Patients with elevated NET levels post-operatively, experienced a fourfold reduction in disease-free survival when compared to the cohorts with lower NET expression (95% CI: 1.39–12.81, p = 0.01). In the same study, using a murine model of liver I/R injury and MC38 cancer cells, the ischaemic group showed increased serum citH3, when compared to controls. After 2 weeks, the liver I/R mice showed greater hepatic metastases compared with mice who did not receive liver I/R injury [26].

The impact of the anaesthetic technique on perioperative NET expression was tested in the first randomised controlled clinical trial in this field on women undergoing breast cancer surgery. NET expression (measured by serum citH3) was assessed in the first randomised control trial (RCT) [68], which evaluated the potential of surgical stress-induced NET formation. The perioperative period represents a window of opportunity for metastasis. The therapeutic options during this phase should not interfere with immunity, wound healing and decrease host-defence [26]. A key aspect of anti-NET therapies is represented by appropriate patient selection. For example, expression of some pro-metastatic factors (e.g. G-CSF) by some tumours and the presence of NETs in those tumours have been proposed as a screening tool to identify patients who may benefit from anti-NET therapies [42]. Although drugs which affect NET formation are clinically available (e.g. domase alpha, lidocaine, PGE-1, etc.), the concept of NET inhibition in cancer is not currently widely supported.

Some potential drugs are experimental (e.g. gasdermin D inhibitor LDC7559), whilst others will have to be relicensed from other pathologies (e.g. HMG-CoA reductase inhibitors, domase alpha, lidocaine). In comparison, analgesics are readily available and may play a double role in the perioperative setting (analgesia/NET inhibition). Of these, cyclooxygenase inhibitors (COX) are the most promising. As the platelet-neutrophil interaction is paramount for NET formation, aspirin (COX inhibitor/antiplatelet agent) has shown promising results, reducing NET production [69]. Although COX blockade leads to prostaglandin generation inhibition, not all COX inhibitors have the same effect on NET expression. In opposition to aspirin, indomethacin (another non-selective COX inhibitor) and diclofenac (COX-2 preferential inhibitor) stimulate NET production by prostaglandin E2 (PGE2) inhibition [70]. In this murine model of transplanted bone marrow neutrophils, indomethacin led to neutrophils’ bactericidal function recovery post-transplant (in vivo) and diclofenac (in vitro), suggesting PGE2 is an inhibitor of NET production [70]. Although various reviews have suggested either an equivocal or slightly positive signal for the use of non-steroidal anti-inflammatory drugs (NSAIDs) in cancer surgery to improve oncological outcomes [33, 71–73], it remains unclear what is the role of NSAIDs in cancer-induced NET expression. The potential NET formation inhibitors are summarised in Table 2.

**NET Formation Modulators**

The perioperative period represents a window of opportunity for metastasis. The therapeutic options during this phase should not interfere with immunity, wound healing and decrease host-defence [26]. A key aspect of anti-NET therapies is represented by appropriate patient selection. For example, expression of some pro-metastatic factors (e.g. G-CSF) by some tumours and the presence of NETs in those tumours have been proposed as a screening tool to identify patients who may benefit from anti-NET therapies [42]. Although drugs which affect NET formation are clinically available (e.g. domase alpha, lidocaine, PGE-1, etc.), the concept of NET inhibition in cancer is not currently widely supported.

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**Conclusion**

Since its discovery, NETs have been shown to be strongly associated with increased cancer progression with potential value as a prognostic indicator and a marker for increased risk of cancer-related pathologies such as metastasis and thrombus formation. The limited observational nature of most of the studies and the paucity of clinical trials preclude a causal link being made. Two recent RCTs investigating the impact of anesthetic technique showed that whilst the addition of lidocaine infusion decreased NET formation, regional anesthesia

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had no effect. Whether modulation of NETs perioperatively translates into improved longer-term oncologic outcomes warrants further evaluation in observational and interventional studies.

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Declarations  

Conflict of Interest  The authors declare no competing interests.

Human and Animal Rights and Informed Consent  This article does not contain any studies with human or animal subjects performed by any of the authors.

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- Of importance
- Of major importance

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