Tumor regionalization after surgery: Roles of the tumor microenvironment and neutrophil extracellular traps

Su-Bin Kwak\textsuperscript{1,2}, Sang Jin Kim\textsuperscript{1,3}, Jiyoung Kim\textsuperscript{1}, Ye-Lim Kang\textsuperscript{1,2}, Chang Woo Ko\textsuperscript{1,2}, Iljin Kim\textsuperscript{4} and Jong-Wan Park\textsuperscript{1,2,3,5} \\
© The Author(s) 2022

Surgery is unanimously regarded as the primary strategy to cure solid tumors in the early stages but is not always used in advanced cases. However, tumor surgery must be carefully considered because the risk of metastasis could be increased by the surgical procedure. Tumor surgery may result in a deep wound, which induces many biological responses favoring tumor metastasis. In particular, NETosis, which is the process of forming neutrophil extracellular traps (NETs), has received attention as a risk factor for surgery-induced metastasis. To reduce cancer mortality, researchers have made efforts to prevent secondary metastasis after resection of the primary tumor. From this point of view, a better understanding of surgery-induced metastasis might provide new strategies for more effective and safer surgical approaches. In this paper, recent insights into the surgical effects on metastasis will be reviewed. Moreover, in-depth opinions about the effects of NETs on metastasis will be discussed.

Experimental & Molecular Medicine (2022) 54:720–729; https://doi.org/10.1038/s12276-022-00784-2

INTRODUCTION
Surgical resection of tumor masses has been regarded as the primary strategy to effectively eradicate cancer cells. Even in cases of incurable cancers, surgery is often performed to prolong the patient’s lifespan or to relieve cancer pain and some complications associated with tumor masses\textsuperscript{1}. However, a wide range of unwanted effects are related to these procedures. In addition to general complications, such as bleeding, thromboembolism, and wound infection, surgery per se can promote cancer metastasis through a series of local and systemic events\textsuperscript{2,3}. A growing body of evidence from clinical and experimental studies has suggested that surgery results in a serious wound that disrupts the structural barrier preventing the outspreading of cancer cells, change the properties of the cancer cells and stromal cells remaining in the tumor microenvironment, or impairs the host defense systems against cancers\textsuperscript{4–6}. Consequently, these unwanted effects of surgery can trigger the second phase of tumor recurrence and metastasis, which are newly acquired events, rather than just outcomes of incomplete treatment. In particular, infection and inflammation during the postoperative period have been reported to increase the risk of cancer recurrence in patients\textsuperscript{6–9}. Notably, these unwanted effects are only found in some patients, not all. Given that the prognostic benefits of surgery are generally greater than the disadvantages, surgical resection should not be abandoned. Nonetheless, it is worthwhile to fully elucidate the mechanisms underlying surgery-triggered aggravation of cancer because it could provide new strategies for more effective and safer surgical approaches for cancers. In this paper, two topics on surgery-triggered cancer metastasis will be discussed. One is a general review concerning the local and systemic influences of cancer surgery on metastasis; the other is the in-depth review concerning the unique role of neutrophils in cancer metastasis.

SURGERY-INDUCED CANCER METASTASIS
Surgeons have long suspected that surgery, even if it is a necessary step in cancer treatment, facilitates cancer metastasis\textsuperscript{3}. This issue remains an unsolved question. Surgery-induced cancer metastasis has been well established in animal models, such as tumor grafts and spontaneous tumorigenesis\textsuperscript{10–12}. As many clinical studies have shown an increased incidence of metastasis during the perioperative period, this event also seems to occur in patients with advanced cancers\textsuperscript{3,13,14}. However, whether the surgical resection of primary tumors is beneficial is controversial. A number of perioperative changes, including tumor cell dissemination, tumor-favoring immune responses, and neoangiogenesis, have been proposed to explain surgery-induced metastasis\textsuperscript{13,14}. This scenario is summarized in Fig. 1.

TUMOR CELL RELEASE BY SURGICAL INJURY
As the tumor cells in the vessels can circulate around the body, they are called circulating tumor cells (CTCs). The number of CTCs is generally viewed as a reliable marker indicating poor outcomes for cancer patients because it is closely associated with tumor metastasis\textsuperscript{15–24}. Moreover, many lines of evidence suggest that CTCs abruptly increase just after surgery\textsuperscript{25–28}. While solid tumors are surgically removed, the tumor architecture is inevitably destroyed due to physical insult, which raises the chance for tumors to shed their cells into the lymphatic and blood vessels.
Even externally palpitating tumors for diagnosis could increase the numbers of CTCs in skin cancer and breast cancer\textsuperscript{27}. However, how significantly surgery-induced CTC release impacts patient prognosis is still controversial. It is not surprising that the CTC number can increase during the perioperative period. However, the CTC number may be dependent on the tumor volume remaining after surgery. It is eventually reduced after complete resection of CTC-releasing tumors but not after incomplete resection\textsuperscript{25}. During surgery, the dissemination and intravasation of tumor cells might be attributed to mechanical pressure and vascular injury\textsuperscript{29}. Long after surgery, tumor cells may spread due to increased cell migration and vascular permeability in the altered microenvironment\textsuperscript{13,30,31}.

**IMMUNITY ALTERED BY SURGICAL STRESS**

Surgical wounds trigger inflammatory responses to favor survival and extravasation of CTCs\textsuperscript{28,32,33}. Surgery can disturb the balance between the innate and adaptive immune systems, leading to impaired surveillance of tumors\textsuperscript{34,35}. Such an effect on surgery seems to be responsible for the systemic response to stress, including the activation of the hypothalamic–pituitary–adrenal (HPA) axis and subsequent production of stress hormones\textsuperscript{36,37}. Upon HPA stimulation, corticotrophin-releasing hormone (CRH) is secreted from the paraventricular nucleus in the hypothalamus and then stimulates the secretion of adrenocorticotropic hormone (ACTH) from the anterior pituitary. The secreted ACTH induces the synthesis and release of glucocorticoids in the adrenal cortex\textsuperscript{38}. As excessive glucocorticoids negatively modulate immune functions, immune surveillance against tumors is considered to be impaired by surgical stress, thereby facilitating the growth of tumors remaining around surgical wounds and metastatic tumors at distant sites\textsuperscript{39–41}.

In addition to glucocorticoids, during stimulation of the HPA axis, the catecholamine hormones epinephrine and norepinephrine are released from the adrenal medulla\textsuperscript{42,43}. Adrenergic receptors (ARs) for these hormones are located on the surface of immune cells.
Many studies have shown that α- and β-ARs are both expressed by innate immune cells, including neutrophils, monocytes, macrophages, dendritic cells, and NK cells. In particular, β2-AR is an AR subtype expressed at the highest level on both innate and adaptive immune cells. Therefore, β2-AR is regarded as the main mediator responsible for the immune effects of catecholamines.

We first review how adrenergic signaling affects innate immunity. β2-AR signaling promotes the M2 differentiation of macrophages, subsequently leading to inhibited production of pro-inflammatory cytokines. In contrast, α-AR signaling has been shown to reverse the effect of β-AR signaling in macrophages. In dendritic cells, β2-AR signaling inhibits their differentiation and antigen presentation function. This signaling also suppresses the production of the pro-inflammatory cytokines IL-12 and TNF-α from dendritic cells, whereas it enhances the production of the anti-inflammatory cytokines IL-6, IL-10, and IL-33. In NK cells, their cytotoxic activity and IFNγ production have been found to be substantially reduced after primary tumors are removed. This NK cell suppression may be attributed to increased levels of catecholamines as well as glucocorticoids. However, the role of adrenergic signaling in the antitumor activity of NK cells is still controversial because it may differ depending on the type and duration of stress. Moreover, there are many reports investigating the adrenergic modulations of other innate immune cells. For example, β2-AR stimulation has been reported to functionally inhibit eosinophils and subsequently aggravate asthma.

In contrast, adrenergic signaling can suppress adaptive immunity. For instance, β-AR signaling decreases the numbers of thymocytes through negative selection in the thymus by activating the p38 signaling pathway. In addition, adrenergic signaling decreases the production of IL-2 and IFN-γ by CD4+T cells and prevents their proliferation. This signaling also regulates Th1 and Th2 differentiation. β2-AR activation was found to promote CD4+ T-cell polarization toward a Th2 phenotype. In memory and effector CD8+ T cells, which express β2-AR at a higher level than naïve T cells, β2-AR signaling downregulates IL-2 and IFN-γ expression under stimulation. In regulatory T cells (Tregs), however, β2-AR signaling enhances immune-suppressive activity by inducing CTLA-4 and Foxp3 expression. The in-depth discussion about this subject is omitted because it is beyond the scope of this review.

ANGIOGENIC STATE ALTERED BY REMOVAL OF PRIMARY TUMORS

A striking hypothesis about angiogenic competition among tumors was experimentally tested by Dr. Folkman and his colleagues. In mice bearing a primary tumor, it was observed that the removal of the primary tumor facilitated the growth of highly vascularized metastases. For hypothetical mechanisms, primary tumors may secrete angiogenic inhibitors as well as angiogenic activators. In the microenvironment of the primary tumor, the activators are abundant enough to overwhelm the inhibitors. In the systemic circulation, however, the activators quickly decay, whereas the inhibitors remain stable. Consequently, small metastases at distant sites are affected to a greater extent by angiogenic inhibitors than by activators, leading to the dormancy of metastases due to limited vascularization. After the primary tumor is surgically removed, the metastases can start to grow vigorously via neoangiogenesis because the circulating inhibitors disappear. Angiostatin and endostatin are regarded as representatives of tumor-derived angiogenesis inhibitors.

ROLES OF NETOSIS IN SURGERY-INDUCED CANCER METASTASIS

Neutrophils are the most abundant type of granulocytes, comprising 40–70% of all white blood cells. These cells neutralize invading microorganisms and act as the main mediators of inflammation. They rapidly accumulate in inflamed areas, where they undergo highly diverse reactions. These cells not only play defensive roles against harmful microorganisms but also clear dead cells for tissue regeneration. However, since neutrophils also play harmful roles in many inflammation-associated diseases, they are considered a double-edged sword. In addition, neutrophils play various roles in the initiation and progression of cancer. In particular, a unique phenomenon named NETosis has been intensively investigated as a pathogenic event in many inflammatory and neoplastic diseases. NETosis refers to the formation of neutrophil extracellular traps (NETs), which are large extracellular complexes composed of chromatin and cytoplasmic/granular proteins. Recently, NETosis has been highlighted as an inflammatory event that promotes cancer metastasis.

NETS AND NETOSIS

Neutrophil extracellular traps (NETs), which were first identified by Volker–Brinkmann and Arturo–Zychlinsky, are fishing net-like structures that can entrap microorganisms invading blood and tissues. Once activated, neutrophils produce intracellular precursors by using DNA, histones, and granular and cytoplasmic proteins and then spread the mature form of NETs out around themselves. A series of these events is called NETosis. In NETs, the following proteins are included: neutrophil elastase, myeloperoxidase, cathepsin G, proteinase 3, lactoferrin, gelatinase, lysozyme, calprotectin, neutrophil defensins, and cathelicidins. As NETs contain a high content of DNA threads, they are sticky enough to entrap and immobilize microorganisms and then kill them using lethal enzymes. Therefore, NETosis is currently defined as an innate immune response against infection. In addition to its antimicrobial activity, NETosis plays a pivotal role in noninfectious autoimmune diseases, such as systemic lupus erythematosus, rheumatoid arthritis, and psoriasis. Moreover, NETosis is involved in other inflammatory diseases, such as vasculitis, intravascular thrombosis, atherosclerosis, periodontitis, and diabetes.

Although the precise mechanisms underlying NETosis are still being investigated, some of the molecular pathways have been identified in two types of NEToses, suicidal (or lytic) NETosis and vital NETosis. Suicidal NETosis mainly depends on the production of reactive oxygen species (ROS). Many stimuli, such as phorbol 12-myristate 13-acetate (PMA), bacterial endotoxins, and interleukins, have been found to induce suicidal NETosis. Indeed, PMA and IL-8 are widely used to induce suicidal NETosis in vitro. Alternatively, suicidal NETosis can be induced by antibodies (especially the Fc region) binding to specific receptors on the surface of neutrophils. Following the stimulation of neutrophils, protein kinase C (PKC) is activated via diverse pathways and in turn activates the Raf-MEK-ERK signaling cascades, followed by ROS production from NADPH oxidases. Alternatively, IL-8 enhances the cytoplasmic levels of ROS, which are mediated by the following process: NF-κB activation via the CXCR2-PJ3-KAK pathway and NOX and COX2 induction by NF-κB. Then, ROS play central roles in suicidal NETosis because they induce the release of the serine protease neutrophil elastase (NE) and myeloperoxidase (MPO) from azurophilic granules and activate peptidyl arginine deiminase 4 (PAD4). PAD4 citrullinates histones in the presence of calcium ions, leading to chromatin decondensation. After the nucleus becomes deformed and ruptured, the decondensed chromatin is released to the cytoplasm and entangled with proteolytic enzymes and other proteins, which are the intracellular precursors of NETs. Finally, the intracellular NET complexes are released through a broken part of the plasma membrane. Since neutrophils die during this process, it is called suicidal NETosis.
However, there is another type of NETosis called vital NETosis. Compared with suicidal NETosis, vital NETosis occurs independently of ROS production from NADPH oxidase. Vital NETosis can be induced by Gram-negative bacteria. LPS in the outer membrane of the bacteria interacts with Toll-like receptor 2 (TLR2) on neutrophils and TLR4 on the surface of platelets, which triggers NETosis. The stimulation of these receptors increases the cytoplasmic level of Ca²⁺ and subsequently stimulates PAD4, leading to NET formation. As NETs are entrapped within vesicles, neutrophils can release them without membrane rupture. The neutrophils undergoing this type of NETosis are still alive, so this process is called vital NETosis. However, the signaling pathway responsible for vital NETosis remains unclear. The molecular mechanisms underlying NETosis are summarized in Fig. 2.

**NETOSIS IN THE TUMOR MICROENVIRONMENT**

Experimental and clinical studies have revealed that NETs are present in a variety of cancers, such as lung cancer, colon cancer, ovarian cancer, and leukemia. This finding suggests that neutrophils actively undergo NETosis in the tumor microenvironment. To date, several hypotheses have been proposed to explain why NETosis is stimulated within tumors. Hypoxia could be one of the reasons because it often develops in growing solid tumors and robustly induces the transcription factor HIF-1. Indeed, McInturff et al. (2012) demonstrated that HIF-1 in neutrophils plays a critical role in NETosis and bacteria-killing activity. Some pro-inflammatory cytokines IL-8, IL-17, G-CSF, CXCL5, and CXCL6 are released from tumor cells and recruit neutrophils in the bone...
marrow to tumor regions. Moreover, many recent reports have suggested that most cytokines potentially initiate or facilitate NETosis in vitro or in vivo. In addition to cytokines, some tumor-derived proteases and tumor exosomes have been reported to induce NETosis. Therefore, NETosis generally actively progresses in the tumor microenvironment.

Recently, NETosis has been investigated as an emerging surrogate marker for cancer diagnosis. The plasma levels of NETs were found to be higher in patients with several types of tumors, including lung cancer, pancreatic cancer, and bladder cancer, than in healthy controls. In lung cancer patients, NETs are present in lung tissues and are also detected in peripheral blood and sputum. In the case of colon cancer, neutrophils from cancer patients were found to undergo NETosis at a higher level under in vitro stimulation than those from healthy controls. Interestingly, such in vitro results positively correlated with the poor clinical outcomes of the patients. Moreover, immunofluorescence staining showed that some components of NETs were detected at the highest level in metastases originating from breast and colon cancers. Recently, a clinical study demonstrated that the serum levels of NET components at the preoperative stage were associated with poor survival in cancer patients. Given these reports, the plasma levels of NET components could be emerging biomarkers to predict the clinical outcomes of cancer patients.

To date, NETosis can be estimated by detecting molecular markers, such as extracellular DNA, citrullinated histone 3 (Cit-H3), and neutrophil-derived protein complexes. However, most researchers tend to confirm this process using more than two markers because each marker has low reliability. Nonetheless, immunochemistry using an anti-Cit-H3 antibody is regarded as a standard method because H3 citrullination is very specific to NETosis. However, even Cit-H3 immunostaining has some drawbacks in use because the procedure is difficult, time-consuming, and not applicable to real-time monitoring. Recently, a research team successfully developed a new NET detection method using the extracellular DNA-intercalating dye CDr15150. This method was found to be faster than previous methods and applicable to the real-time tracing of NETosis. Considering the clinical importance of NETosis, it is worthwhile to develop new NET-detecting materials that are applicable to experimental and clinical studies.

NEUTROPHIL ACTIVATION AND NETOSIS FOLLOWING SURGICAL INJURY

Surgical trauma and subsequent complications, including wound infection, increase neutrophil counts in the peripheral blood through increased granulopoiesis in the bone marrow. Additionally, tissue injury affects the function of circulating neutrophils. The changes in neutrophil functions depend on the micromilieu of the damaged tissue. In wounds, damaged and necrotic cells express the signals responsible for early neutrophil recruitment, which are damage-associated molecular patterns (DAMPs). DAMPs are cellular components including DNA, histones, ATP, interleukin-1α, high mobility group protein B1 (HMGB1), N-formyl peptides, and others. DAMPs directly activate neutrophils through G-protein-coupled receptors. Indirectly, DAMPs can stimulate surrounding tissues to produce chemokines and lipid mediators for neutrophil chemotaxis. In wounds, neutrophils clear necrotic cells and invading microorganisms. Moreover, they release various cytokines responsible for tissue repair. Indeed, neutrophils express and store a variety of growth factors and angiogenic factors that contribute to regeneration and revascularization. For example, the proliferative cytokines TGFβ and IL-10 and the angiogenic factor VEGF are representative of neutrophil-derived tissue repair proteins.

As mentioned above, NETosis is a defense system to protect the body from invading pathogens. However, when neutrophils are excessively stimulated, they produce excess NETs, thereby leading to pathological consequences. To eliminate a visceral tumor, surgeons make a deep incision on the skin, remove a part of the tumor-bearing organ, and reconstruct surrounding tissues for functional recovery. Since such a procedure involves large wounds, it can induce severe inflammation and subsequently stimulate neutrophils to undergo NETosis. Given many clinical reports showing that the plasma levels of NETosis markers are elevated after major surgeries, NETs formed at surgical wounds are believed to circulate throughout the body along with blood and lymphatic streams. Since NETs circulate in the peripheral blood, the components of NETs could be surrogate markers for evaluating the progression of cancers in clinical settings.

ROLES OF NETS IN TUMOR METASTASIS

Metastasis refers to tumor cell spread out of the original location and the formation of secondary lesions at distant sites. Tumor cells metastasize through the following sequences: local invasion, intravasation into the blood or lymphatic vessels, escape from the immune system, anchoring to capillaries in target organs, extravasation into the organs, transformation from dormant cells to proliferating cells, colonization to micrometastases, and growth to macrometastases. Many lines of evidence suggest that NETs promote metastasis at multiple steps, which are summarized in Fig. 3. NETs promote the local invasion of cancer cells by degrading the extracellular matrix (ECM). ECMs, which are mainly composed of fibrous proteins and polysaccharides, act as a barrier blocking cell movement, and thus, degrading the ECM is essential for tumor cell invasion. As mentioned previously, NETs contain many proteolytic enzymes, such as neutrophil elastase, matrix metalloproteinase 9, and cathepsin G. These NET enzymes loosen the ECMs to allow cancer cell movement. NETs also promote the intravasation of cancer cells by increasing vascular permeability. The wall of a capillary consists of two layers: the endothelial layer and the outer basement membrane (mainly made of connective tissues). Capillaries (especially, continuous capillaries) are characterized by a complete endothelial lining with tight junctions between endothelial cells. The tight junction is impermeable to macromolecules but allows the passage of small molecules, such as water, ions, gases, metabolites, and hormones. Therefore, cancer cells should penetrate the two layers to enter the circulation, which cannot occur normally. Given that the basement membrane and the tight junction are both mainly composed of proteins, it is not surprising that protease-containing NETs loosen the vascular wall and allow cancer cells to penetrate into the vessel. Moreover, pro-inflammatory cytokines released from activated neutrophils escalate the enhancement of vascular permeability.

Even if cancer cells successfully enter the circulation, only a few of them can survive to finally establish metastases. Indeed, millions of tumor cells are released into the circulation every day, but metastases are not observed as frequently as we estimate. There are two obstacles that circulating tumor cells (CTCs) need to overcome: the shearing force by blood flow and the immune system. NETs can wrap up CTCs with platelets owing to their sticky properties. In lung cancer patients, neutrophils from cancer patients were found to undergo NETosis at a higher level under in vitro stimulation than those from healthy controls. Interestingly, such in vitro results positively correlated with the poor clinical outcomes of the patients. Moreover, immunofluorescence staining showed that some components of NETs were detected at the highest level in metastases originating from breast and colon cancers. Recently, a clinical study demonstrated that the serum levels of NET components at the preoperative stage were associated with poor survival in cancer patients. Given these reports, the plasma levels of NET components could be emerging biomarkers to predict the clinical outcomes of cancer patients.

To date, NETosis can be estimated by detecting molecular markers, such as extracellular DNA, citrullinated histone 3 (Cit-H3), and neutrophil-derived protein complexes. However, most researchers tend to confirm this process using more than two markers because each marker has low reliability. Nonetheless, immunochemistry using an anti-Cit-H3 antibody is regarded as a standard method because H3 citrullination is very specific to NETosis. However, even Cit-H3 immunostaining has some drawbacks in use because the procedure is difficult, time-consuming, and not applicable to real-time monitoring. Recently, a research team successfully developed a new NET detection method using the extracellular DNA-intercalating dye CDr15150. This method was found to be faster than previous methods and applicable to the real-time tracing of NETosis. Considering the clinical importance of NETosis, it is worthwhile to develop new NET-detecting materials that are applicable to experimental and clinical studies.
this journey may be a new focus of metastasis. In the growth of micrometastases, the involvement of NETs has been demonstrated in several animal studies. For instance, when tumor-bearing mice were injected with DNase (decomposing NETs), the growth of the preexisting metastases was significantly retarded. In mice subjected to abdominal surgery, NETs in the peritoneum collected tumor cells and provided a microenvironment favoring tumor survival and growth. These results suggest that NETosis is a potential target to prevent surgery-induced tumor metastasis.

After metastasizing to distant tissues, tumor cells are often found to remain dormant for a period of time and unexpectedly regrow later. To date, little is known about the molecular mechanisms underlying tumor dormancy and reactivation. In addition to the metastatic process, NETs are believed to participate in the reactivation of dormant cancer cells in metastatic regions. An animal study showed that dormant micrometastases became aggressively growing metastases after lung inflammation was induced by tobacco smoke or nasal instillation of lipopolysaccharide. In this study, the NET-associated proteases NE and MMP-9 were found to be responsible for the reactivation of dormant cancer cells. Laminin cleavage by enzymes may induce reactivation through the integrin α3β1 signaling pathway. However, the effects of NETs on the reactivation of dormant cells remain uncertain.

CONCLUSION
Here, we discuss how distant metastasis paradoxically increases after resection of the primary tumor. From two points of view, this topic was discussed: one is a general review concerning surgery-triggered metastasis; the other is a review concerning the roles of NETs in metastasis. After surgical removal of tumors, the tumors and surrounding tissues are harshly handled, which increases the chance for tumor cells to spread. Because the surgery results in a deep wound, it might provoke systemic stress, which inhibits immunity against tumor cells. Moreover, as small metastases can escape from the angiogenic control of the primary tumor, they can grow vigorously after tumor surgery and are clinically detected as multiple metastases. Second, we discuss recent advances in tumor- and surgery-induced NETosis. NETosis is an innate immune process that entraps and kills microorganisms, but it also plays a pathogenic role in many inflammatory diseases. Neutrophils are activated and subsequently undergo NETosis in the tumor microenvironment and the surgical wound, both of which are enriched with pro-inflammatory cytokines. Furthermore, many reports have suggested that NETs stimulate the entire metastatic process from local invasion of cancer cells to colonization/growth. Therefore, NETosis could be an emerging target for blocking tumor metastasis after tumor surgery.

In fact, surgery is a very complicated procedure because it is accompanied by many medications for anesthesia, analgesia, muscle relaxation, and infection control. Obviously, these medications during the perioperative period should be considered risk factors that may affect tumor metastasis. Nonetheless, we restrictively focused on the effects of surgical procedures and wounds on tumor cell spreading and metastasizing to distant organs. Considering that surgery-related medications are doctors’ most likely options, we can try to substitute safer drugs for...
metastasis-inducing drugs. We hope that this topic will be discussed in future studies.

REFERENCES

1. Wyld, L., Audisio, R. A. & Poston, G. J. The evolution of cancer surgery and future perspectives. Nat. Rev. Clin. Oncol. 12, 115–124 (2015).
2. Demichelis, R., Retzky, M. W., Hrushesky, W. J., Baum, M. & Gukas, I. D. The effects of surgery on tumor growth: a century of investigations. Ann. Oncol. 19, 1821–1828 (2008).
3. Tohme, S., Simmons, R. L. & Tsung, A. Surgery for cancer: a trigger for metastases. Cancer Res. 77, 1548–1552 (2017).
4. Alieva, M., van Rheejen, J. & Broekman, M. L. D. Potential impact of invasive surgical procedures on primary tumor growth and metastasis. Clin. Exp. Metastasis. 35, 319–331 (2018).
5. Chen, Z. et al. Surgical stress and cancer progression: the twisted mango. Mol. Cancer https://doi.org/10.1186/s12943-019-1058-3 (2019).
6. Tang, F., Tie, Y., Tu, C. & Wei, X. Surgical trauma-induced immunosuppression in cancer: recent advances and the potential therapies. Clin. Transl. Med. 10, 199–223 (2020).
7. Nojiri, T. et al. Long-term impact of postoperative complications on cancer recurrence following lung cancer surgery. Ann. Surg. Oncol. 24, 1135–1142 (2017).
8. Matsubara, D. et al. The impact of postoperative complications on recurrence in patients with colorectal cancer. Int. J. Clin. Oncol. 25, 602–610 (2020).
9. Tsujimoto, H. et al. Impact of postoperative infectious complications on adjuvant chemotherapy administration after gastrectomy for advanced gastric cancer. Jpn. J. Clin. Oncol. 51, 379–386 (2021).
10. Zhang, Y., Zhang, N., Hoffman, R. M. & Zhao, M. Surgically-induced multi-organ metastasis in an orthotopic syngeneic imageable model of 4T1 murine breast cancer. Anticancer Res. 35, 4641–4646 (2015).
11. Tsuchiya, Y. et al. Increased surgical stress promotes tumor metastasis. Surgery 133, 547–555 (2003).
12. Al-Sahaf, O., Wang, J. H., Browne, T. J., Cotter, T. G. & Redmond, H. P. Surgical injury enhances the expression of genes that mediate breast cancer metastasis to the lung. Ann. Surg. 252, 1037–1043 (2010).
13. Alieva, M. et al. Preventing inflammation inhibits biopsy-mediated changes in tumor cell behavior. Sci. Rep. https://doi.org/10.1038/s41598-017-07660-4 (2017).
14. Paterlini-Brechot, P. & Benali, N. L. Circulating tumor cells (CTC) detection: recent advances and the potential therapies. Langenbecks Arch. Surg. 390, 333–341 (2005).
15. Lin, E., Calvano, S. E. & Lowry, S. F. Inflammatory cytokines and cell response in surgery. Surgery 127, 117–126 (2000).
16. Gibbison, B., Angelini, G. D. & Lightman, S. L. Dynamic output and control of the hypothalamic-pituitary-adrenal axis in critical illness and major surgery. Br. J. Anaesth. 111, 347–360 (2013).
17. Dietzey, F. G., Leker, R. R. & Weidenfeld, J. Activation of the adrenocortical axis by surgical stress: involvement of central norepinephrine and interleukin-1. Neuroimmunomodulation 7, 182–188 (2000).
18. Gibbison, B. et al. Dynamic pituitary-adrenal interactions in response to cardiac surgery. Crit. Care Med. 33, 791–800 (2015).
19. Brokaw, J. I. et al. Glucocorticoid-induced apoptosis of dendritic cells in the rat tracheal mucosa. Am. J. Respir. Cell Mol. Biol. 19, 598–605 (1998).
20. Piemonti, L. et al. Glucocorticoids affect human dendritic cell differentiation and maturation. J. Immunol. 162, 4673–4681 (1999).
21. Rea, D. et al. Glucocorticoids transform CD40-triggering of dendritic cells into an alternative activation pathway resulting in antigen-presenting cells that secrete IL-10. Blood 95, 3162–3167 (2000).
22. McCoy, L. K. Physiology of the autonomic nervous system. Am. J. Pharm. Educ. https://doi.org/10.5688/aj710478 (2007).
23. Sun, N. et al. High-resolution tissue mass spectrometry imaging reflects a refined functional anatomy of the human adult adrenal gland. Endocrinology 159, 1511–1524 (2018).
24. Nance, D. M. & Sanders, V. M. Autonomic innervation and regulation of the immune system (1987–2007). Brain Behav. Immun. 21, 736–745 (2007).
25. Ordovas-Montanes, J. et al. The regulation of immunological processes by peripheral neurons in homeostasis and disease. Trends Immunol. 36, 578–604 (2015).
26. Cosentino, M. et al. Endogenous catecholamine synthesis, metabolism, storage and uptake in human neurons. Life Sci. 64, 975–981 (1999).
27. Elenkov, I. J., Wilder, R. L., Chrousos, G. P. & Vizi, E. S. The sympathetic nerve–an integrative interface between two subsystems: the brain and the immune system. Pharmacol. Rev. 52, 595–638 (2000).
28. Straub, R. H. Complexity of the bi-directional neuroimmune junction in the spleen. Trends Pharmacol. Sci. 25, 640–646 (2004).
29. Herve, J. et al. β2-Adrenoceptor agonist inhibits antigen cross-presentation by dendritic cells. J. Immunol. 190, 3163–3171 (2013).
30. Sanders, V. M. The beta2-adrenergic receptor on T and B lymphocytes: do we understand it yet? Brain Behav. Immun. 26, 195–202 (2012).
31. Maestroni, G. J. & Mazzola, P. Langerhans cells β2-adrenoceptors: role in migration, cytokine production, Th priming and contact hypersensitivity. J. Neuroimmunol. 144, 91–99 (2003).
32. Salicru, A. N. et al. Variable effects of combination corticosteroids and catecholamines on total beta 2-adrenergic receptors in human CD4+ T-lymphocytes. J. Allergy Clin. Immun. https://doi.org/10.1016/j.jaci.2006.11.585 (2007).
33. Tang, L. et al. Sympathetic nerve activity maintains an anti-inflammatory state in adipose tissue in male mice by inhibiting TNF-alpha gene expression in macrophages. Endocrinology 156, 3680–3694 (2015).
34. Liu, B. et al. Leptin-mediated increases in catecholamine signaling reduce adipose tissue inflammation via activation of macrophage HDA4. Cell Metab. 19, 1058–1065 (2014).
35. Grailer, J. J., Haggadone, M. D., Sarma, J. V., Zeteoune, F. S. & Ward, P. A. Induction of M2 regulatory macrophages through the β2-adrenergic receptor with protection during endotoxemia and acute lung injury. J. Innate Immun. 6, 607–618 (2014).
56. Garcia, J. L., del Carmen Saez, M., De la Fuente, M. & Ortega. Regulation of phagocytic process of macrophages by noradrenaline and its end metabolite 4-hydroxy-3-metoxophenyl-glycol. Role of α- and β-adrenoceptors. Mol. Cell. Biochem. 254, 299–304 (2003).

57. Qin, J. F. et al. Adrenergic receptor β2 activation by stress promotes breast cancer progression through macrophages M2 polarization in tumor microenvironment. BMC Cancer 18, 289–300 (2018).

58. Sloan, E. K. et al. The sympathetic nervous system induces a metastatic switch in primary breast cancer. Cancer Res. 70, 7042–7052 (2010).

59. Agac, D., Estrada, L. D., Maples, R., Hooper, L. V. & Farrar, J. D. The β2-adrenergic receptor controls inflammation by driving rapid IL-10 secretion. Brain Behav. Immun. 74, 176–185 (2018).

60. Chen, C. et al. β2-adrenergic receptors stimulate interleukin-6 production through Epac-dependent activation of PKCdelta/p38 MAPK signalling in neonatal mouse cardiac fibroblasts. Br. J. Pharmacol. 166, 676–688 (2012).

61. Yanagawa, Y., Matsumoto, M. & Togashi, H. Adrenergoc-uppermented enhancement of interleukin-33 production by dendritic cells. Brain Behav. Immun. 25, 1427–1433 (2011).

62. Anglìa, L. et al. Natural killer cell IFNα expression is profoundly suppressed following colorectal cancer surgery. Ann. Surg. Oncol. 25, 3747–3754 (2018).

63. Reinhardt, R. et al. Invasive surgery impairs the regulatory function of human CD56 bright natural killer cells in response to Staphylococcus aureus. Suppression of interferon-gamma synthesis. PLoS One https://doi.org/10.1371/journal.pone.0130155 (2015).

64. Nair, M. P. & Schwartz, S. A. Immunomodulatory effects of corticosteroids on breast cancer growth. Cancer Res. 70, 7042–7052 (2010).

65. Hahnfeldt, P., Panigrahy, D., Folkman, J. & Hlatky, L. Tumor development under angiogenic signaling: a dynamical theory of tumor growth, treatment response, and postvascular dormancy. Cancer Res. 59, 4770–4775 (1999).

66. O'Reilly, M. S. et al. Endostatin: an endogenous inhibitor of angiogenesis and tumor growth. Cell 88, 277–285 (1997).

67. Holmgren, L., O'Reilly, M. S. & Folkman, J. Dormancy of micrometastases: balanced proliferation and apoptosis in the presence of angiogenesis suppression. Nat. Med. 1, 149–153 (1995).

68. Phillips, M. & Kubel, P. The neutrophil in vascular inflammation. Nat. Med. 17, 1331–1339 (2011).

69. Ley, K., Laudanna, C., Cybulsky, M. I. & Nourshargh, S. Getting to the site of inflammation: the leukocyte adhesion cascade updated. Nat. Rev. Immunol. 7, 678–689 (2007).

70. Brinkmann, V. et al. Neutrophil extracellular traps kill bacteria. Science 303, 1532–1535 (2004).

71. Mantovani, A., Cassatellia, M. A., Costantini, C. & Jaillon, S. Neutrophils in the activation and regulation of innate and adaptive immunity. Nat. Rev. Immunol. 11, 519–531 (2011).

72. guarantees, http://www.neuroendoology.com Neutrophil extracellular traps promote the development of reparative macrophages mediated by ROS to orchestrate liver repair. Nat. Commun. https://doi. org/10.1038/s41467-019-09046-8 (2019).

73. Kuhl, A. A. et al. Aggravation of different types of experimental colitis by depletion or adhesion blockade of neutrophils. Gastroenterology 133, 1882–1889 (2007).

74. Headland, S. E. et al. Neutrophil-derived microvesicles enter cartilage and protect the joint in inflammatory arthritis. Sci. Transl. Med. 7, 315ra190 (2015).

75. Parkos, C. A. Neutrophil-epithelial interactions: a two-edged sword. Am. J. Pathol. 186, 1404–1416 (2016).

76. Tazawa, H. et al. Infiltration of neutrophils is required for acquisition of metastatic phenotype of benign murine fibrosarcoma cells: implication of inflammation-associated carcinogenesis and tumor progression. Am. J. Pathol. 163, 2221–2232 (2003).

77. Lokinouva, E. et al. Growth regulated oncogene-alpha expression by murine squamous cell carcinoma promotes tumor growth, metastasis, leukocyte infiltration and angiogenesis. Exp. Cell Res. 240, 1–9 (2010).

78. Sangaletti, S. et al. Neutrophil extracellular traps mediate transfer of cytoplasmic neutrophil antigens to myeloid dendritic cells toward ANCA induction and associated autoimmunity. Blood 120, 3007–3018 (2012).

79. Hu, S. C. et al. Neutrophil extracellular trap formation is increased in psoriasis and induces human beta-defensin-2 production in epidermal keratinocytes. Sci. Rep. https://doi.org/10.1038/srep11119 (2016).

80. Guglietta, S. et al. Coagulation induced by C3aR-dependent NETosis drives protumorigenic neutrophils during small intestinal tumorigenesis. Nat. Commun. https://doi.org/10.1038/s41467-019-11037-6 (2017).

81. Garcia-Romo, G. S. et al. Netting neutrophils are major inducers of type I IFN production in pediatric systemic lupus erythematosus. Sci. Transl. Med. 3, 73ra20 (2011).

82. Garcia-Romo, G. S. et al. Neutrophil extracellular traps mediate transfer of cytoplasmic neutrophil antigens to myeloid dendritic cells toward ANCA induction and associated autoimmunity. Blood 120, 3007–3018 (2012).

83. Hu, S. C. et al. Neutrophil extracellular trap formation is increased in psoriasis and induces human beta-defensin-2 production in epidermal keratinocytes. Sci. Rep. https://doi.org/10.1038/srep11119 (2016).

84. Guglietta, S. et al. Coagulation induced by C3aR-dependent NETosis drives protumorigenic neutrophils during small intestinal tumorigenesis. Nat. Commun. https://doi.org/10.1038/s41467-019-11037-6 (2017).

85. Rispens, T. Neutrophil extracellular traps accumulate in peripheral blood and compromise organ function in tumor-bearing animals. Cancer Res. 75, 2653–2662 (2015).

86. Fuchs, T. A. et al. Novel cell death program leads to neutrophil extracellular traps. J. Cell Biol. 176, 231–241 (2007).

87. Brinkmann, V. & Zychlinsky, A. Beneficial suicide: why neutrophils die to make NETs. Nat. Rev. Microbiol. 5, 577–582 (2007).

88. Masucci, M. T., Minopoli, M., Del Vecchio, S. & Carrierio, M. V. The emerging role of neutrophil extracellular traps (NETs) in tumor progression and metastasis. Front. Immunol. https://doi.org/10.3389/fimmu.2020.01749 (2020).

89. Snoeck, H. T., Boone, B. A. & Bennewitz, M. F. Neutrophil extracellular traps in breast cancer and beyond: current perspectives on NET stimuli, thrombosis and metastasis, and clinical utility for diagnosis and treatment. Breast Cancer Res. https://doi.org/10.1186/s13058-019-1237-6 (2019).

90. Korkmaz, B. Moreau, T. & Gauthier, F. Neutrophil elastase, protease 3 and cathepsin G: physicochemical properties, activity and physiopathological functions. Biochimie 90, 227–232 (2008).

91. Neumann, A. et al. The antimicrobial peptide LL-37 facilitates the formation of neutrophil extracellular traps. Biochem. J. 464, 3–11 (2014).

92. Papayannopoulos, V., Metzler, K. D., Hakkim, A. & Zychlinsky, A. Neutrophil elastase and myeloperoxidase regulate the formation of neutrophil extracellular traps. J. Cell Biol. 191, 677–691 (2010).
Urban, C. F. et al. Neutrophil extracellular traps contain calprotectin, a cytosolic protein complex involved in host defense against Candida albicans. Plos Pathog. https://doi.org/10.1371/journal.ppat.1000639 (2009).

Urban, C. F., Reichard, U., Brinkmann, V. & Zychlinsky, A. Neutrophil extracellular traps capture and kill Candida albicans yeast and hyphal forms. Cell. Microbiol. 8, 668–676 (2006).

Villanueva, E. et al. Netting neutrophils induce endothelial damage, infiltrate tissues, and expose immunostimulatory molecules in systemic lupus erythematosus. J. Immunol. 187, 538–552 (2011).

Soderberg, D. et al. Increased levels of neutrophil extracellular trap remnants in the circulation of patients with small vessel vasculitis, but an inverse correlation to anti-neutrophil cytoplasmic antibodies during remission. Rheumatology 54, 2085–2094 (2015).

Martinod, K. et al. Neutrophil elastase-deficient mice form neutrophil extracellular traps in an experimental model of deep vein thrombosis. J. Thromb. Haemost. 14, 551–558 (2016).

Nahrendorf, M. & Swirski, F. K. Neutrophil-macrophage communication in inflammation and atherosclerosis. Science 349, 237–238 (2015).

Vilkov, L., Klappacher, M., Hannig, M. & Kräutgartner, W. D. Extracellular neutrophil traps in periodontitis. J. Periodontal Res. 44, 664–672 (2009).

Wong, S. L. et al. Diabetes primes neutrophils to undergo NETosis, which impairs wound healing. Nat. Med. 21, 815–819 (2015).

Ypp, B. G. & Kubes, P. NETosis: how vital is it? Blood 122, 2784–2794 (2013).

Behnhen, M. et al. Immobilized immune complexes induce neutrophil extracellular trap release by human neutrophil granulocytes via Fc gamma RIIB and the Fc receptor. J. Immunol. 193, 1954–1965 (2014).

Gray, R. D. et al. Activation of conventional protein kinase C (PKC) is critical in the generation of human neutrophil extracellular traps. J. Inflamm. https://doi.org/10.1186/1476-9255-10-12 (2013).

Hakkim, A. et al. Activation of the Raf-MEK-ERK pathway is required for neutrophil extracellular trap formation via induction of hypoxia-inducible factor 1 alpha. J. Exp. Med. 207, 1217–1225 (2010).

Shijo, H. et al. Evaluation of neutrophil functions after experimental abdominal sepsis. J. Pathol. 213, 463–474 (2016).

Huang, C. & Niethammer, P. Tissue damage signaling is a prerequisite for pro-inflammation and neutrophil extracellular trap formation: Implications for the establishment of cancer-associated thrombosis. Sci. Rep. https://doi.org/10.1038/s41598-017-06893-7 (2017).

Orlova, V. V. et al. A novel pathway of HMGB1-mediated inflammation and neutrophil extracellular trap production in patients with colorectal cancer in vitro. Inflamm. Res. https://doi.org/10.1155/2017/4915062 (2016).

Barbu, E. A., Mendelsohn, L., Samsel, L. & Thein, S. L. Pro-inflammatory cytokines associate with NETosis during sickle cell vaso-occlusive crises. Cytokine https://doi.org/10.1016/j.cytokine.2019.154933 (2020).

Visser, T., Hietbrink, F., Groeneveld, K. M., Koenderman, L. & Leenen, L. P. Isolated neutrophil extracellular traps are increased in cancer patients but does not associate with venous thrombosis. Cardiovasc. Diagn. Ther. 7, S140–S149 (2017).

Rosinski, M., Yarmush, M. L. & Berthiaume, F. Quantitative dynamics of in vivo neutrophil extracellular traps: Implications for the establishment of cancer-associated thrombosis. Blood 132, 1109–1120 (2018).

Boettcher, S. et al. Endothelial cells translate pathogen signals into G-CSF-driven emergency granulopoiesis. Blood 124, 1393–1403 (2014).

Zhang, Q. et al. Circulating mitochondrial DAMPs cause inflammatory responses to injury. Nature 464, 104–107 (2010).

Visser, T., Hietbrink, F., Groeneveld, K. M., Koenderman, L. & Leenen, L. P. Isolated blunt chest injury leads to transient activation of circulating neutrophils. Eur. J. Trauma Emer. Surg. 33, 676 (2007).

Richards, J. R. R., Hendrickse, C., Gao-Smith, F. & Thickett, D. R. Neutrophil extracellular trap production is upregulated in patients with colorectal cancer in vitro. J. Inflamm. https://doi.org/10.1186/1749-7763-10-1120 (2013).

Oltowy, O. O., Zhu, X., Cass, D. L. & Smith, C. W. Neutrophil recruitment by fetal porcine endothelial cells: implications in scarless fetal wound healing. Pediatr. Res. 55, 464–469 (2007).

McInturf, A. M. et al. Mammalian target of rapamycin regulates neutrophil extracellular trap formation via induction of hypoxia-inducible factor 1 alpha. Blood 120, 3118–3125 (2012).

Verbeke, H. et al. Isotopic neutralizing antibodies against mouse GCP-2/CXCL6 inhibit melanoma growth and metastasis. Cancer Lett. 302, 54–62 (2011).

Demers, M. et al. Priming of neutrophils toward NETosis promotes tumor growth. Oncoimmunology https://doi.org/10.2147/Oimm20190354 (2020).

Xu, X. et al. CXCL5 promotes gastric cancer metastasis by inducing epithelial-mesenchymal transition and activating neutrophils. Oncogenesis https://doi.org/10.1038/s41399-020-00249-z (2020).

Gupta, A. K., Hasler, P., Holzgrewe, W., Gebhardt, S. & Hahn, S. Induction of neutrophil extracellular DNA lattices by placental microparticles and IL-8 and their presence in plasma. Hum. Immunol. 66, 1146–1154 (2005).

Gupta, A. K. et al. Activated endothelial cells induce neutrophil extracellular traps and are susceptible to NETosis-mediated cell death. FEBS Lett. 584, 3193–3197 (2010).

Zhang, Y. et al. Interleukin-17-induced neutrophil extracellular traps mediate resistance to checkpoint blockade in pancreatic cancer. J. Exp. Med. https://doi.org/10.1084/jem.20190354 (2020).
167. Lazaro-Diez, M. et al. Human neutrophils phagocytose and kill Acinetobacter baumannii and A. pittii. Sci. Rep. https://doi.org/10.1038/s41598-017-04870-8 (2017).

168. Simpson, D. M. & Ross, R. The neutrophilic leukocyte in wound repair a study with antineutrophil serum. J. Clin. Invest. 51, 2009–2023 (1972).

169. Nishio, N., Okawa, Y., Sakurai, H. & Isobe, K. Neutrophil depletion delays wound repair in aged mice. Age 30, 11–19 (2008).

170. Tecchio, C. & Cassatella, M. A. Neutrophil-derived cytokines involved in physiological and pathological angiogenesis. Chem. Immunol. Allergy 99, 123–137 (2014).

171. Aldabbous, L. et al. Neutrophil extracellular traps promote angiogenesis: evidence from vascular pathology in pulmonary hypertension. Arterioscler. Thromb. Vasc. Biol. 36, 2078–2087 (2016).

172. Yipp, B. G. et al. Infection-induced NETosis is a dynamic process involving neutrophil multiassembling in vivo. Nat. Med. 18, 1386–1393 (2012).

173. van Dam, L. S., Rabelink, T. J., van Kooten, C. & Teng, Y. K. O. Clinical implications of excessive neutrophil extracellular trap formation in renal autoimmune diseases. Kidney Int. Rep. 4, 196–211 (2019).

174. Stakos, D., Skendros, P., Konstantinides, S. & Ritits, K. Traps N'clots: NET-mediated thrombosis and related diseases. Thromb. Haemost. 120, 373–383 (2020).

175. von Meijenfeldt, F. A. et al. Elevated plasma levels of cell-free DNA during liver transplantation are associated with activation of coagulation. Liver Transplant 24, 1716–1725 (2018).

176. Paunel-Gorgulu, A. et al. cfDNA correlates with endothelial damage after cardiac surgery with prolonged cardiopulmonary bypass and amplifies NETosis in an intracellular TLR9-independent manner. Sci. Rep. https://doi.org/10.1038/s41598-017-17561-1 (2017).

177. Banki, F. et al. Plasma DNA as a molecular marker for completeness of resection and recurrent disease in patients with esophageal cancer. Arch. Surg. Clin. 142, 533–538 (2007).

178. Ludovini, V. et al. Plasma DNA, microsatellite alterations, and p53 tumor mutations are associated with disease-free survival in radically resected non-small cell lung cancer patients: a study of the perugia multidisciplinary team for thoracic oncology. J. Thorac. Oncol. 3, 365–373 (2008).

179. Deckers, A. S. et al. Prognostic role of blood NETosis in the progression of head and neck cancer. Cells https://doi.org/10.3390/cells8090946 (2019).

180. Park, J. et al. Cancer cells induce metastasis-supporting neutrophil extracellular DNA traps. Sci. Transl. Med. 8, 361ra138 (2016).

181. Thalini, C. et al. Citrullinated histone H3 as a novel prognostic blood marker in patients with advanced cancer. Plos One https://doi.org/10.1371/journal.pone.0109123 (2018).

182. Khan, U. et al. Neutrophil extracellular traps in colorectal cancer progression and metastasis. Int. J. Mol. Sci. https://doi.org/10.3390/ijms22147260 (2021).

183. Estrella, V. et al. Acidity generated by the tumor microenvironment drives local invasion. Cancer Res. 73, 1524–1535 (2013).

184. Seliger, B. Strategies of tumor immune evasion. BioDrug 19, 347–354 (2005).

185. Zavyalova, M. V. et al. Intravasation as a key step in cancer metastasis. Biochemistry 84, 762–772 (2019).

186. Desgroiseller, J. S. et al. An integrin alphavbeta3(C)-Src oncopgenic unit promotes anchorage-independence and tumor progression. Nat. Med. 15, 1163–1169 (2009).

187. Strilic, B. & Offermann, S. Intravascular survival and extravasation of tumor cells. Cancer cell 32, 282–293 (2017).

188. Phan, T. G. & Croucher, P. I. The dormant cancer cell life cycle. Nat. Rev. Cancer 20, 398–411 (2020).

189. Parhi, L. et al. Breast cancer colonization by Fusobacterium nucleatum accelerates tumor growth and metastatic progression. Nat. Commun. https://doi.org/10.1038/s41467-020-16967-2 (2020).

190. Coolen-Lartigue, J., Spicer, J., Najmeh, S. & Ferri, L. Neutrophil extracellular traps in cancer progression. Cell. Mol. Life Sci. 71, 4179–4194 (2014).

191. Das, A., Monteiro, M., Barai, A., Kumar, S. & Sen, S. MMP proteolytic activity regulates cancer invasiveness by modulating integrins. Sci. Rep. https://doi.org/10.1038/s41598-017-14340-w (2017).

192. Liotta, L. A. Adhere, degrade, and move: The three-step model of invasion. Cancer Res. 76, 3115–3117 (2016).

193. Deryugina, E. et al. Neutrophil elastase facilitates tumor cell intravasation and early metastatic events. iScience https://doi.org/10.1016/j.isci.2020.101799 (2020).

194. Bjork, J., Hedqvist, P. & Arfors, K. E. Increase in vascular permeability induced by leukotriene B4 and the role of polymorphonuclear leukocytes. Inflammation 6, 189–200 (1982).

195. Coolen-Lartigue, J. et al. Neutrophil extracellular traps sequester circulating tumor cells and promote metastasis. J. Clin. Invest. 123, 3446–3458 (2013).

196. Najmeh, S. et al. Neutrophil extracellular traps sequester circulating tumor cells via beta1-integrin mediated interactions. Int. J. Cancer 140, 2321–2330 (2017).

197. Park, S. Y. & Nam, J. S. The force awakens: metastatic dormant cancer cells. Exp. Mol. Med. 52, 569–581 (2020).

198. Albrengues, J. et al. Neutrophil extracellular traps produced during inflammation awaken dormant cancer cells in mice. Science https://doi.org/10.1126/science.aao4227 (2018).

ACKNOWLEDGEMENTS
J.-W.P. was supported by the Education and Research Encouragement Fund of Seoul National University Hospital.

AUTHOR CONTRIBUTIONS
J.-W.P. collected the information and wrote the article. S.B.K. designed and drew the images. S.-J.K., J.K., Y.L.K., C.W.K., and I.K. provided substantial contributions to the discussion of the content. All authors reviewed the manuscript before submission. This work was supported by grants from the National Research Foundation of Korea (NRF-2019R1A2B5B03069677).

COMPETING INTERESTS
The authors declare no competing interests.

ADDITIONAL INFORMATION
Correspondence and requests for materials should be addressed to Jong-Wan Park.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

© The Author(s) 2022