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IL-12 induced modulation of tumor derived myeloid cell inflammatory responsiveness.

Courtney Jetun Mitchell
University of Louisville

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IL-12 INDUCED MODULATION OF TUMOR DERIVED MYELOID CELL INFLAMMATORY RESPONSIVENESS

By

Courtney Jetun Mitchell
B.S. Clark Atlanta University, 2007
M.D. University of Louisville, 2014

A Dissertation
Submitted to the Faculty of the
University of Louisville School of Medicine
in Partial Fulfillment of the Requirements
for the Degree of

Doctor of Philosophy

Department of Microbiology and Immunology
University of Louisville
Louisville, KY

August, 2014
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A Dissertation Approved on
May 23, 2014

by the following Dissertation Committee:

_________________________________
Jill Suttles

_________________________________
Haval Shirwan

_________________________________
Nejat Egilmez

_________________________________
Jason Chesney
DEDICATION

Dedicated to my beloved mother, Jacqueline Marie Green (1964-2007).
ACKNOWLEDGMENTS

Many have been instrumental in getting me to this point. First, I would like to thank the late Dr. Robert “Bob” Stout. Bob was such a great mentor. No matter what time of day or what else he had going on as chair of the department; he always had an open door for me. He was always ready to engage and discuss my next great idea, no matter how far-flung it might have been. He cared deeply about the students in the department, and supported me whole-heartedly during my project. Even at the close of his life, he was concerned with my path to getting my PhD. He challenged my thinking, and provided an example of strength under tremendous suffering. I will always be indebted to him for all he taught me and his unwavering support of all my research ambitions. Thank you Bob. I hope that this work is a reflection of the great man and scientist you were.

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Finally, I praise God for his continued blessings and the opportunities He has allowed in my life.
Tumor infiltrating and tumor associated myeloid cells (TIMs and TAMs) elaborate an array of factors that promote tumor growth and metastasis. IL-12, a potent inflammatory cytokine has been shown to induce regression of many cancers. We hypothesize that IL-12 augments the ability of TIMs and TAMs to respond to inflammatory stimuli providing a window in which these stimuli are more likely to promote tumor destruction. Related to this hypothesis, we asked two questions: (1) Does IL-12 directly change signaling events associated with inflammatory signal transduction? (2) Is IFNγ required for the entirety of IL-12 induced enhancement of the response of TIMs to inflammatory stimuli? First, we looked broadly at the in vivo effects of microspheres containing IL-12 on the growth and metastasis of 4T1 tumors. We also did studies with tumor cell-dendritic cell fusion for future examination of the impact of IL-12 and myeloid cells on the efficacy of tumor cell-dendritic cell fusion vaccines in mice. Most of this dissertation
focuses on in vitro work using TIMs and TAMs isolated from wild-type BALB/c or IFNγ deficient mice bearing the 4T1 mammary carcinoma. TIMs and TAMs were pretreated in vitro with IL-12 followed by LPS. TNFα, IL-6, and IL-10 cytokine and mRNA levels were measured. We also examined the impact of IL-12 on the response 4T1 TIMs to tumor derived products. The phosphorylation of a number molecules involved in inflammatory signaling pathways, including MAPK proteins, were assessed by Western Blot. We found that treatment of TIMs with IL-12 followed by exposure to LPS enhances the amount of IL-6 and TNFα with a reciprocal decrease in IL-10. This observation is associated with increases in the phosphorylation of MAPKs. The presence of IFNγ is only partially necessary for these effects. We observed that IL-12 only significantly impacted the production of IL-10 from 4T1 TAMs in response to LPS. IL-12 caused a significant increase in the amount of TNFα and IL-6 in response to tumor derived products without affecting IL-10. Our results provide additional insight into direct changes induced by IL-12 to the functional phenotype of TIMs and TAMs in response to inflammatory stimuli.
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CHAPTER I
INTRODUCTION

Myeloid Derived Cell Phenotypic and Functional Diversity

Myeloid derived cells are heterogeneous innate immune cells from the myeloid lineage of bone marrow-derived hematopoietic stem cell precursors. These cells include monocytes, macrophages, neutrophils, dendritic cells, and myeloid derived suppressor cells. Macrophages are phagocytic cells broadly distributed in tissues within the body, and are mature forms of circulating blood monocytes (1). These cells are critical for innate immune responses and assist in priming the adaptive immune system to attack pathological insults (1).

Macrophages are remarkably dynamic and display an array of functions and activities. In addition to their phagocytic properties, macrophages are capable of presenting antigen to adaptive immune cells, secreting factors important for wound healing, and establishing the extracellular matrix (1). Interestingly, many macrophage functions are in direct opposition to one another. Inflammation provides an excellent illustration of the diverse functions of macrophages. In the formative stages of inflammation, macrophages secrete pro-inflammatory cytokines, chemokines as well as cytotoxic factors that activate surrounding immune and stromal cells, attract auxiliary innate immune cells and specialized adaptive immune cells. All of these functions work
together to initiate the killing or clearance of the offending inflammatory agent (2, 3). As inflammation progresses, macrophages secrete less pro-inflammatory cytokines and begin to elaborate factors that aid in the resolution of inflammation, wound healing, and angiogenesis (2, 3).

As evidence of the spectrum of functions macrophages may exhibit, several subsets of macrophages have been described in the literature. Discussions continue as to whether these subsets are distinct lineages of myeloid cells locked into a particular functional state or transient cellular phenotypes that change given the stimulus or the environment (2-5). A simplistic classification of macrophages based on T-helper cell categorization was introduced by Mills, labeling macrophages as either M1 or classically activated and M2 or alternatively activated (6). Macrophage secretion of pro-inflammatory cytokines, such as TNF-alpha (TNF$\alpha$), IL-1 beta (IL-1$\beta$), IL-12, IL-6, IL-15, metalloproteinases (MMPs) and nitric oxide, and expression of pro-inflammatory genes, such as iNOS and co-stimulatory molecules B7 and major histocompatability class II molecules (MHC II) in response to IFN gamma (IFN$\gamma$) followed by LPS treatment, generally characterize classically activated macrophages (7, 8). Conversely, macrophages exposed to IL-13 in combination with IL-4 display the alternative activated phenotype characterized by the secretion of anti-inflammatory cytokines like IL-10, transforming growth factor-beta (TGF$\beta$), vascular endothelial growth factor (VEGF) and the expression of anti-inflammatory genes including, arginase I and mannose receptor I (9). Attempts to incorporate the complex and diverse functional phenotypes of
Macrophages have been made by the use of sub-classifications of the original M1-M2 paradigm (7) or a color wheel for macrophage classification based on the functional environment, the major examples being immune regulation, wound healing, and host defense (3, 10, 11). Overall, macrophages are remarkably diverse cells that display a plethora of functional phenotypes under the influence of various environmental factors.

**Macrophage Activation and Counter-Regulation**

*Activation*

Macrophages can be induced to display various functional phenotypes by many activating agents. Upon encountering various signals in the environment, macrophages can be activated to secrete cytokines and mediators of inflammation, present antigens for adaptive immune cell activation, and kill intracellular and extracellular bacteria. Specific examples of macrophage activators include bacterial lipopolysaccharide (LPS), CD40 ligand (CD40L or CD154), and IFNγ.

The activation of macrophages via LPS has been extensively studied. Macrophage stimulation with LPS leads to a variety of effector functions through the integration of many complex intracellular signaling pathways. LPS is a component of the outer membrane of gram-negative bacteria (12). LPS acts as an endotoxin and binds to the pattern recognition receptor (PRR), toll-like receptor 4 (TLR4) and co-receptors CD14 and MD-2 on macrophages (13-15). Intracellular signals from the LPS/TLR4/MD-2 complex are transmitted via
adaptor proteins, myeloid differentiation factor 88 (MyD88) or TIR-containing adaptor molecule (TRIF) (16). Early TLR4 signaling via the MyD88 pathway is responsible for elaborating most of the pro-inflammatory cytokines classically associated with LPS activation of macrophages (17). The MyD88 pathway leads to the degradation of inhibitors of kappa beta (IκBs), which allows for the activation and nuclear translocation of NF-κB (18). NF-κB is a protein complex consisting of dimers of any of its 5 family members that include p65, p50, Rel, A, Rel B, and c-Rel (19). The dimerization of the p65 and p50 subunits is associated with LPS stimulation (20). The delayed MyD88 independent, TRIF-biased pathway signals through tumor necrosis factor receptor-associated factor (TRAF) family member associated NF-κB-activator binding kinase (TBK1) to activate IFN response factor 3 (IRF3) (21-23). This leads to the release of type I interferons such as interferon B (IFNβ) (21). The TRIF-biased pathway can also lead to the delayed activation of NF-κB (23)

The CD40 signaling pathway in macrophages has also been thoroughly investigated (24). CD40 is a member of the tumor necrosis factor receptor (TNFR) family (25, 26). This co-stimulatory molecule is found on the surface of macrophages, B-cells, T cells, dendritic cells, and endothelial cells (27-30). Engagement of CD40 on macrophages, by its ligand, CD154 leads to secretion of pro-inflammatory cytokines and elaboration of reactive oxygen species (31-33). CD154 is a member of the TNF family of proteins (34). It can be found bound to the cell surface of various immune cells or in a soluble form (34). Introduction of CD154 in tumor bearing mice has been shown to decrease the
number of suppressive myeloid cells and increase T cell activation (34). Signaling events downstream of CD40 include the phosphorylation of protein tyrosine kinases (PTKs), such as Janus activated kinase 3 (JAK3) (35), which can lead to the activation of transcription factor signal transducer and activator of transcription 3 (STAT3) and STAT6 (35, 36). CD40 ligation has been shown to lead to activation of serine/threonine kinases: c-jun amino-terminal kinase/stress activated protein kinase (JNK/SAPK), p38 mitogen-activated protein kinase (MAPK), and extracellular signal-regulated mitogen-activated kinase (ERK) (37, 38). Activation of CD40 by CD154 also enlists tumor necrosis factor receptor-associated factors (TRAFs) to transmit signaling cascades to downstream transcriptional mediators (39). TRAF2, TRAF3, TRAF5, and TRAF6 have shown to play a role in CD40 signaling (40). CD40 signaling can result in the activation of NF-κB via TRAF2 and TRAF6, and the activator protein 1 (AP-1), and nuclear factor of activated T cells (NF-AT), all of which can mediate inflammatory cytokine production (40-43).

Another example of a macrophage activator is IFNγ. IFNγ is the sole member of the type II interferon group (44). Th1 lymphocytes, NK cells, and macrophages produce IFNγ, which is a potent endogenous activator of macrophages (45-48). Genes that are upregulated in macrophages in response to IFNγ include MHC class I and II, anti-viral proteins, inflammatory cytokines and chemokines (46, 49). IFNγ binds to its receptor to form a heterodimer of type II cytokine receptors, interferon gamma receptor gamma 1 and 2 (IFNGR1 and 2)
that leads to the activation of JAK1 and JAK2 that subsequently phosphorylate IFNGR1 (50, 51). This phosphorylation event provides a docking site for STAT1 (50-52). STAT1 phosphorylation allows it to dimerize and translocate into the nucleus to bind to target genes on interferon response elements (52, 53). IFNγ has been shown to prime macrophages to be tumoricidal and for subsequent stimulation by activating stimuli such as LPS (54-56). Overall, macrophages are activated by a variety of stimuli that integrate numerous signaling pathways leading to diverse functional phenotypes.

Counter-Regulation

Of equal importance to macrophage activation is macrophage counter-regulation. Excessive inflammation can cause tissue damage and pathology. Thus, the dampening and resolution of macrophage inflammatory activities are very important. The immune system has evolved to include several mechanisms to negatively regulate inflammation. Macrophage activation can be attenuated in several ways.

In addition to the pro-inflammatory cytokines secreted by macrophages in response to LPS, concurrent signaling events lead to the elaboration of IL-10, a potent anti-inflammatory cytokine. Twenty years ago, it was first demonstrated that IL-10 was produced by human monocytes in response to LPS (57). The same study showed that IL-10 inhibited the release of IL-1α and β, IL-6, TNFα, and IL-8 in response to LPS and/or IFNγ (57). Subsequent reports have unmasked a complex network of activators and signaling components that
regulate IL-10 expression in macrophages and mediate its downstream anti-inflammatory effects. Several transcriptional factors have been implicated in the regulation of IL-10 in macrophages. These include specificity factor (Sp) activators, STATs, c-Musculoaponeurotic Fibrosarcoma (c-MAF), activator proteins (AP-1), CCAAT/Enhancer binding proteins (C/EBP), and cAMP response element binding protein (CREB) (58). Sp1 binds to the G-rich portion of the IL-10 promoter (59, 60). Lack of endogenous Sp protein leads to complete ablation of IL-10 promoter activity (59). STAT1 and STAT3 recruitment to the IL-10 promoter has been demonstrated, although evidence of direct binding of STAT proteins to existing cis-elements within the IL-10 promoter is not available (61, 62). In macrophages, the protooncogene, c-MAF, a member of the basic leucine zipper family of transcription factors is constitutively present and is ushered to the IL-10 promoter upon TLR ligation in response to LPS (63). C/EBPs can act as homo- and heterodimers in TLR-dependent and independent pathways to impact IL-10 gene expression (58). In response to LPS, C/EBP proteins synergize with Sp1 to enhance IL-10 transcription (64). Alternatively, cAMP stimulation through the adenosine receptor can lead to activation of the IL-10 promoter in a MyD88-independent mechanism via C/EBP (65). MyD88-independent activation of the IL-10 promoter may also be due to CREB activation (66). In response to cAMP stimulation, CREB is phosphorylated and accompanied into the nucleus with its co-activators, histone acetyl transferases, CREB binding protein (CBP) and p300(67, 68). This leads to hyperacetylation of histones at the IL-10 promoter and ultimately, increased transcription of IL-10
Phosphorylation of serine 10 on histone H3 is thought to be a critical epigenetic marker for transcriptional activation of the IL-10 gene (69).

IL-10 binds to its receptor, IL-10 receptor (IL-10R), a tetramer consisting of two heterodimers of the two subunits, IL-10R1 and IL-10R2 (70). Upon ligation, IL-10R1 associates with JAK1 that is then phosphorylated (71). JAK1 phosphorylation and activation leads to the phosphorylation of STAT3, an indispensable mediator of IL-10 anti-inflammatory function (71). STAT3 enters the nucleus and facilitates the transcription of several anti-inflammatory genes including ETS family transcriptional repressor, ETV3, a helicase family corepressor, Strawberry notch homologue 2 (SBNO2), and suppressor of cytokine signaling 3 (SOCS3) (72). SOCS3 is part of an 8-member family of proteins characterized by a Src homology 2 (SH2) domain, a variable N-terminus and a conserved region termed the SOCS box at its C-terminus. Two members of this group, including SOCS3 have a kinase inhibitory region (KIR) at the N-terminal (73). SOCS3 binds phosphorylated tyrosines on cytokine receptors, especially the IL-6 receptor component gp130, to inhibit signal transduction. SOCS3 is thought to be key molecule in distinguishing between IL-10 and IL-6 cellular signaling (74, 75). Although both IL-6 and IL-10 induce STAT3 and SOCS3, SOCS3 selective inhibition of the IL-6 receptor component, gp130, disallows IL-6 from displaying anti-inflammatory characteristics associated with IL-10 (74, 75). A STAT3 independent pathway for IL-10 gene expression has been elucidated that includes phosphatidylinositol 3’ kinase (PI3K), glycogen synthase kinase 3 beta (GSK3β), and Akt, which controls IL-10 induced
expression of autotaxin, Egr2, caspase5, and signaling lymphatic activation molecule (SLAM) (76).

Because persistent high levels of IL-10 lead to immunosuppression and render the host susceptible to pathogenic insult, negative regulation of IL-10 is of vital importance. IL-10 induced SOCS3 can inhibit STAT3 activity creating a negative autoregulatory loop (75). Additionally, MHC class II transactivator (CIITA), and poly (ADP) ribose polymerase 1 (PARP-1) (in an allelic specific manner) have been shown to negatively regulate IL-10 expression in innate immune cells (77, 78). Interestingly, studies have demonstrated that IFNγ can inhibit IL-10 production in response to LPS (79, 80). It was shown that IFNγ inhibits IL-10 through diminishing the effects of the MAPK pathway, and the PI3K/Akt/GSK3b pathways (80).

The MAPK signaling pathway

Optimal intracellular communication in macrophages and other immune cells requires a complex system of highly integrated and tightly controlled signals. Various signaling pathways must converge upon numerous transcription factors and ultimately genes to regulate the function of macrophages. These same pathways may exert post-transcriptional and post-translational modifications to further dictate the macrophage functional phenotype, adding another layer of complexity to the signaling tableau. The MAPK pathway is an example of a single signaling pathway that can influence a broad spectrum of other signaling pathways. This pathway has been shown to be involved in a wide
variety of signaling cascades that control myeloid cell response to inflammatory stimuli. The expansive influence of the MAPK pathway on the macrophage functional phenotype is evident as this pathway modulates and contributes to the propagation of signals transmitted by LPS, IFNγ, and CD40.

MAPKs are family of proteins that form a tiered system for relaying signals within cells (81). In response to a stimulus, MAPK kinase kinases (MAP3Ks) are activated. These activated MAP3Ks go on to activate MAPK kinases (MAP2Ks or MEKs) via phosphorylation of specific serine and threonine residues. Finally, MAP2Ks proceed to activate MAPK family members, ERKs, p38s, and JNKs, through phosphorylation of specific tyrosines and threonines. Once activated MAPKs can bind and activate various transcription factors and enhancers to regulate gene transcription (81).

The first MAPKs discovered were ERK family members including ERK1, ERK2, MAPK1, and MAPK3 (82). ERK family members are encoded by two genes, ERK1 and ERK2 and require dual phosphorylation at Thr203 and Tyr205 (ERK1) and Thr185 and Tyr187 (ERK2) for activation (82). ERKs can be activated by a well-defined pathway initiated by the Ras protooncogene, or via Ras-independent pro-inflammatory stimuli such as TNFα and LPS (83).

JNKs were first described as protein kinases found in the livers of cycloheximide treated rats (83). Like ERKs, JNKs have a variety of activators. JNKs are extremely sensitive to mitogens, environmental stressors like heat shock and ionizing radiation, pro-inflammatory cytokines, endoplasmic reticular (ER) stress, DAMPs, and PAMPs (81). There are 3 JNK family members, JNK1-3
Genes of the same name encode them, with each family member having several isoforms as a result of alternative splicing. JNKs are dually phosphorylated at Thr 183 and Tyr185 (82). A large number of MAP3Ks have been shown to lead to the activation of JNKs including MEKK1, MEKK2, MLK1, MLK2, and MLK4 (84). These MAP3Ks lead to the phosphorylation of two MAP2Ks, MKK4 and MKK7. Both MKK4 and MKK7 are required for JNK phosphorylation (85, 86). A well-known substrate for JNKs is AP-1.

A third MAPK family member is p38. There are four p38 genes and identically named proteins: $p38\alpha$ (MAPK14), $p38\beta$ (MAPK11), $p38\gamma$ (MAPK12), and $p38\delta$ (MAPK13) (83). The $p38\alpha$ and $p38\beta$ isoforms are ubiquitously expressed, while the other isoforms are distributed with more tissue specificity (82). Most literature refers to $p38\alpha$ because it is the most abundant of the two ubiquitous forms. Like the two previously mentioned MAPKs, p38 is activated by a wide variety of PAMPs, DAMPs, pro-inflammatory cytokines, and environmental stressors. Activation of p38 requires phosphorylation at Thr180 and Tyr182 (82).

Given the wide distribution and extensive activating signals of the MAPKs, it is not unexpected that it has been shown that these proteins are critical to the macrophage’s response to stimuli. In response to LPS TLR4 ligation, ERK1/2, JNK, and p38 are all activated by both the early MyD88 dependent pathways and late MyD88 independent, TRIF dependent routes (82). LPS triggers the activation of the MAP3K, tumor locus progression 2 (Tpl2 aka Cot3 and MAP3K8). Tpl2 activates MKK1 (MAP2K1) and MKK2 (MAP2K2) (87). This leads to ERK1/2
activation (87). This pathway has been shown to be crucial and required for LPS induced TNFα and prostaglandin E2 (PGE2) secretion (87-89).

Of the three MAPK family members, p38 has been demonstrated to be the most critical MAPK mediator of LPS provoked gene transcription (90). Illustrating the importance of p38α in the inflammatory phenotype of macrophages, conditional deletion of p38α in macrophages significantly decreased the secretion of pro-inflammatory cytokines, TNFα, IL-12, and IL-18 in response to LPS in vitro, and diminished sepsis in two in vivo animal models (91). This is apparently due to the extreme reduction in activation of CREB and CEB/P when p38−/− macrophages are exposed to LPS (91). It is thought the MAPK activated protein kinase 2 (MK2) is the downstream target of p38 that controls CREB activation (92). This pathway is also important for IL-10 and COX2 production (92).

The MAPK family proteins are also implicated in the IFNγ signaling pathway. Like LPS, IFNγ induces p38, ERK1/2, and JNK (93, 94). It was shown that p38 is involved with controlling the level of expression of genes required for the innate immune response, including CCL5, CXCL9, and CXCL10; cytokines such as TNF-α; and inducible NO synthase (iNOS). Alternatively, JNK-1 was shown to affect genes involved in antigen presentation like CIITA and MHC class II molecules (93).

**Myeloid Cells and Cancer**
As demonstrated by the intricate integration of multiple signaling pathways, understanding the continuously changing macrophage and other myeloid cells functional phenotype under normal physiological conditions is a complex proposition alone. In a diverse and complicated pathological condition such as cancer, deciphering the function and role of macrophages and myeloid cells presents a unique challenge. The idea of inflammation being connected with the initiation of cancer can be traced back to Virchow’s 1863 observation of the “lymphoreticular infiltrate” in cancer and his postulation that cancer may arise at sites of chronic inflammation (95). Conversely, the importance of the immune system in defending against cancer has also been appreciated. Activation of the immune system with “Coley’s Toxin,” a mixture of attenuated bacterial products, was shown to induce the regression of osteosarcoma (96). Later studies have shown that immunization of mice with irradiated tumor cells prevents the growth of the same viable tumor cells but not different, unrelated, viable tumor cells (97). The evidence supporting the influence of the immune system has on initiation, growth, spread, and eradication of cancer prompted Weinberg et al., to list “avoiding immune destruction” and “tumor promoting inflammation” as emerging hallmarks in his latest update of the highly cited Hallmarks of Cancer: The Next Generation (98, 99).

Macrophages and other myeloid-derived cells have been shown to be critical for several aspects of tumor initiation, progression, and metastasis. Several groups have shown that the presence of high numbers of myeloid cells within tumors is associated with a poor prognosis in many cancers (100-103).
Chemically induced, obesity-driven and virus-mediated cancer models provide clear links between inflammation and cancer initiation. A report showing that the anti-inflammatory protein SOCS3 decreases dextran sodium sulfate (DSS) and azoxymethane (AOM) associated tumorigenesis in the gut highlights the role of inflammation in chemically induced cancers (104). A recent study showed that human adipose tissue macrophages (ATMs) induced lipid accumulation and an inflammatory phenotype in human cancer cell lines (105). Additionally, these ATMs displayed genes associated with cancer related pathways and were genetically similar to TAMs (105). Macrophages were also found to be associated with malignant transformation of the human papilloma virus-infected cervix (106). Once tumors are established, macrophages and other myeloid cells including myeloid-derived suppressor cells (MDSCs) display an immunosuppressive, anti-inflammatory phenotype that assists tumor cells in evading immune detection. MDSCs are a heterogeneous subset of immature myeloid cells that can be grouped as either granulocytic (CD11b+, Ly6G+) or monocytic (CD11b+, Ly6C+) (107-109). It is reasonable to speculate that these cells are a product of the remarkable plasticity of macrophages and myeloid cells and the ability of macrophages to respond to various environmental factors (2). C-C chemokine receptor 2+ (CCR2+), CD11b+ monocytic MDSCs were shown to aid in tumor immune escape by limiting the infiltration of activated T cells into the tumor (110). One study revealed that TAMs in renal cell carcinoma secrete high amounts of chemokine (C-C) ligand 2 (CCL2) and IL-10 that contribute to
immunosuppression and perpetuate the recruitment and maintenance of more TAMs (111).

The Pollard et al., was one of the first to definitively show that macrophages were vital for cancer metastatic progression when they demonstrated that mice lacking macrophage colony stimulating factor-1 (CSF-1) did not have changes in tumor incidence or growth, but had delayed development of invasive, metastatic cancers (112). This study highlighted one of the most critical roles macrophages perform over the course of a cancer. The ability of macrophages to promote tumor invasiveness and metastasis through modulating stromal elements and supporting abnormal vascularization and angiogenesis has subsequently been thoroughly researched. Hypoxia within the tumor microenvironment causes TAMs to secrete VEGF that supports immature blood vessel formation leading to vascular leakiness, which promotes cancer metastasis (113). A study identified a subset of TAMs that express the angiopoietin receptor, TIE2 that is critical for macrophage induced angiogenesis (114). Also important for tumor spread is degradation of the surrounding stroma and extracellular matrix. Macrophages have been shown to secrete high amounts of MMPs that can degrade tissue and allow metastasis (115, 116).

In addition to tumor infiltrating myeloid cells and macrophages found within the tumor that work to suppress the immune system and aid in tumor escape from the original site, several studies have shown that TAMs away from the primary tumor work to prime distal locations for cancer metastasis. It has
been demonstrated that macrophages are essential for metastasis and growth of tumor cells at distant sites (112). Additionally, it has been illustrated that macrophages in the lung are key in initial stages of cancer cell seeding and extravasation (117). Qian et al., demonstrated that distal tissue stromal cells and early metastatic cancer cells secrete CCL2 that attracts inflammatory monocytes and macrophages (118). This study showed that these macrophages promote the extravasation and growth of tumor cells at distant sites, in part, via the secretion of VEGF (118). Additionally, it was recently reported that tissue factor mediated coagulation helped recruit macrophages to pre-metastatic niches and this recruitment is necessary for metastatic cancer cell survival (119). It has also been shown that macrophages support tumor cell survival at distal sites by expressing α4-integrin on their surface which binds to cancer cell expressed vascular cell adhesion molecule 1 (VCAM-1) (116). This α4-integrin-VCAM-1 interaction leads to the activation of pro-survival signals in the cancer cells via the phosphoinositide 3-kinase (PI3K)/AKT pathway (116).

The dichotomous role of macrophages and myeloid cells in cancer, with their ability to initiate and promote tumor growth, while suppressing their own capability to thwart cancer progression, provide an opportunity for targeting these cells in anti-cancer therapies.

**Myeloid Cells and Immunotherapy**

*Current Immunotherapeutic Approaches and Challenges*
In addition to the traditional therapeutic approaches to cancer including chemotherapy, radiation therapy, and surgery, immunotherapy has reemerged as a viable component of a successful cancer treatment plan. The recent FDA approval of Sipuleucel-T (Provenge), an autologous immunotherapeutic, illustrates the potential impact of thoughtful and effective immunotherapeutic agents (120, 121). Sipuleucel-T treatment requires that a patient’s peripheral mononuclear blood cells (PMBC) be removed and exposed to a fusion protein consisting of prostatic acid phosphatase (PAP) and the cytokine, GM-CSF (120, 121). This treatment results in both clinical and statistical improvement of prostate cancer patient outcomes (120, 121). Another instance in which the effectiveness of immunotherapy is evident is in the use of Ipilimumab (Yervoy), a human monoclonal antibody that antagonizes a negative regulator of cytotoxic T cells, cytotoxic T-lymphocyte antigen 4 (CTLA-4), to render cytotoxic T cells more active against melanoma (122, 123).

It is generally accepted that a major difficulty in cancer immunotherapy is the robust immunosuppressive environment established in part by myeloid cells. An approach to alleviating this profound immunosuppression is the exploitation of the functional plasticity of myeloid cells. As demonstrated by earlier cited literature, macrophages and other myeloid cells have the remarkable ability to respond to environmental signals including the tumor microenvironment. The strategy of altering the phenotype of macrophages from a tumor and metastasis-supportive, immunosuppressive phenotype to an anti-tumorigenic, pro-inflammatory phenotype would provide an avenue to activate the immune system.
to eradicate cancers. One such agent that has been shown to have anti-tumor effects and changes the functional profile of macrophages is the cytokine IL-12.

**The Biology of IL-12**

Michiko Kobayashi and Giorgio Trinchieri first described IL-12 in 1989 as NK cell stimulatory factor (NKSF) (124). IL-12 is a heterodimeric cytokine composed of two disulfide-linked subunits, p35 and p40 that together form the p70 unit (125, 126). The p35 subunit shares homology with other cytokines such as granulocyte colony stimulating factor (G-CSF) and IL-6, while the p40 heavy chain is shared with IL-23 and is a member of the hemapoietin receptor family along with ciliary neurotrophic factor (CNTF) and the IL-6 receptor (127). IL-12p40 can be secreted as a monomer or a homodimer and is thought to have antagonistic activity, while IL-12p35 is not found in a monomeric form (128). The genes for the IL-12 subunits are located on separate chromosomes. In humans p35 is on chromosome 3 and p40 is on chromosome 5, while in mice p35 and p40 are on chromosomes 6 and 11, respectively (125). IL-12 is secreted by a variety of immune cells including monocytes and macrophages, neutrophils, dendritic cells, and B cells (129). IL-12 can initiate pro-inflammatory functions in NK cells, T cells, and myeloid cells, making it an attractive, multivalent agent to reverse the strongly immunosuppressive tumor microenvironment (129). IL-12 is a strong inducer of IFNγ from T cells, NK cells, and myeloid cells; it also induces immune cell proliferation, and enhances cell-mediated toxicity (129).
The *IL-12p40* gene is regulated at the transcriptional level (130). The *p40* gene can be induced by a variety of microbial products. The p40 promoter has been shown to have binding sites for several transcription factors including NF-κB, IRF-1, c-Rel, interferon consensus binding protein, and Ets family members (130-134). Deletion of the cis-elements at the Ets or NF-κB binding site restricts *p40* promoter activity (135). Conversely to p40, p35 is ubiquitously expressed and controlled both transcriptionally and translationally (136). Although there is constitutive synthesis of the p35 message in unstimulated cells, scant protein is actually produced due to an inhibitory ATG sequence at the 5' untranslated region (136). When a stimulus is applied to cells, the transcriptional start site is altered, and the inhibitory region is not included in the message (136).

Many DAMPs and PAMPs can prompt IL-12 production. They include LPS, Lipoteichoic acid (LTA), peptidoglycan, and bacterial (CpG) DNA, and they work independent of T cells (137, 138). IL-12 is also made in response to antigen presenting cell CD40 ligation by CD40L on T cells (139). A positive feedback loop is established when IL-12 induced IFNγ triggers more IL-12 production in a macrophage cell line (131, 140). Alternatively, IL-12 synthesis is attenuated by IL-10, IL-11, IL-13, and type I IFNs (141-143). It has also been demonstrated in macrophages that stimulation of G-coupled protein receptors (GCPRs) including the receptors for MCP-1, prostaglandin E2, histamine, and Fc receptor activation can inhibit IL-12 production (144-147). Interestingly, cholera toxin, an activator of a GPCR subunit, and the measles virus via CD46 inhibit IL-12 production (148-151).
IL-12 binds to its receptor, IL-12 receptor (IL-12R)(152). The IL-12R consists of 2 chains, β1 and β2 (152). They are members of the type I cytokine receptor family with close homology to IL-6 receptor component, gp130 (127). The human *IL-12β1* and *IL-12β2* genes are found on chromosomes 19 and 1 in humans and on chromosomes 8 and 6 in mice, respectively. The IL-12 receptor is expressed on NK cells, T cells, dendritic cells, and macrophages. Co-expression of both chains confers optimal responsiveness to IL-12 although the β2-subunit is often used to predict IL-12 activity due to the presence of three cytoplasmic tyrosine residues on the β2-chain (127, 153).

The IL-12 signal is transmitted from the IL-12R via the JAK-STAT pathway (127). The β1-receptor chain interacts with JAK2 while the β2-receptor subunit associates with the tyrosine kinase, TYK2 (154, 155). The previously mentioned tyrosine residues on the β2-receptor are phosphorylated upon IL-12 ligation, which allows for the recruitment, phosphorylation, and activation of STAT4. Although it has also been reported that STAT1 and STAT3 are activated by IL-12, STAT4 is thought to be the major component of the IL-12 signaling pathway as STAT4 deficient mice display an identical phenotype to IL-12p40 deficient mice (156, 157). STAT4 binds to both the human and murine *Ifng* gene (157-159). IL-12 stimulation also results in the phosphorylation of Src family tyrosine kinase, Lck and the MAPK, p38 (160-162). Activation of p38 by IL-12 appears to be important for IFNγ production and Th1 differentiation (162). STAT4 has been shown to be negatively regulated by the ubiquitin E3 ligase, SLIM (163).
Much of IL-12’s physiological influence has been attributed to its activation of adaptive components of the immune system and NK cells and its elicitation of IFNγ. IL-12 can directly induce the proliferation of pre-activated T and NK cells and augment the proliferation of resting and naïve cells caused by several mitogenic agents (153, 164). IL-12 can also enhance the generation of cytotoxic T lymphocytes (CTL), NK and lymphokine activated (LAK) cells by eliciting the transcription of genes such as those for granzymes, perforin, and adhesion molecules that render these cells more cytotoxic (164). One of the essential roles of IL-12 is the induction of IFNγ secretion by T cells, B cells, and NK cells (153, 165). IL-12 alone, or in synergy with IL-18, a member of the IL-1 cytokine family can cause IFNγ secretion (166-168). The IL-12-IFNγ axis in conjunction with signaling through the T cell receptor (TCR) and other receptors is crucial for Th1 differentiation (153).

**IL-12 and Myeloid Cells**

More attention has been placed on how IL-12 may be involved with other immune cells including myeloid cells. Most of this work has focused on the autocrine effects of IFNγ from IL-12 stimulated macrophages. Macrophages have been shown to produce nitric oxide and upregulate surface molecules associated with antigen presentation in response to IL-12 (169). Both of these effects are dependent on IFNγ as this phenotype is erased in IFNγ-/- mice (165, 170). Additionally, IL-12 has been demonstrated to prime macrophages to produce copious amounts of NO in response to LPS and IFNγ (171). Some work also
suggests that IL-12 may regulate TGF-β1 production since macrophages from mice deficient in p40 chain of IL-12 secrete high amounts of TGF-β1 (172).

**IL-12 as a Cancer Immunotherapeutic**

Because IL-12 can activate both the innate and adaptive arms of the immune system, much effort has been used to investigate its effectiveness as an anti-tumor agent and therapeutic vaccine adjuvant. Initial studies in several animal models of cancer yielded promising results regarding IL-12 alone or in combination with other cytokines as an immunotherapeutic drug and vaccine adjuvant. IL-12 has been shown to be very effective in eradicating tumors in mouse models of breast cancer, melanoma, thymoma, renal cell carcinoma, colon cancer, and lung cancer (173-178). However, early clinical trials with IL-12 were unsuccessful, largely due to the profound cytotoxicity associated with systemic administration of IL-12 (179). Researchers continue to work to find safe alternatives for administering IL-12 in humans. There are currently several active clinical trials involving IL-12 and various cancers. These phase I and phase II clinical trials include IL-12 in combination with a dendritic cell breast cancer fusion vaccine (NCT00622401), the heavy chain of the antibody of NHS576 which targets double-stranded and single-stranded DNA that are exposed in areas of necrosis (NCT01417546), a plasmid DNA vaccine for Merkel cell cancer (NCT0144081816), the monoclonal antibodies cetuximab and daclizumab (NCT01468896 and NCT01307618), and T cells engineered to secrete IL-12 in metastatic melanoma (NCT012365718) (clinicaltrials.gov). Most of IL-12's in vivo
anti-tumor effect has been attributed to CD8+ and CD4+ T cells, as well as NK cells. It has also been reported that IL-12 can activate NKp46+ lymphoid tissue-inducer cells (LTi cells) to induce tumor rejection (177). Recently the impact of IL-12 on TAMs has been studied. Our lab showed that IL-12 modified the functional phenotype of TAMs in vivo and in vitro and that TAMs released cytoplasmic IL-15 in response to IL-12 in vivo, and that this event was critical for the initiation of IL-12’s anti-tumor activity (173, 180). It was more recently shown that IL-12 enhances the ability of myeloid cells within tumors to improve the activation and cytotoxicity of CD8+ T cells (175).
Rationale, Significance, and Hypothesis

Our attention to the relationship between IL-12 and tumor associated macrophages an myeloid cells is well-founded given the accumulating evidence showing myeloid cells as remarkably plastic, the significant influence these cells have on cancer growth, metastasis, and immunosuppression, and the use of IL-12 as an anti-cancer therapy. Understanding the impact of IL-12 on TIMs and TAMs is an important component in deciphering the mechanisms related to IL-12 immunotherapeutic effects.

Recent work has identified TIMs and TAMs as direct targets of IL-12 (175). What is not completely clear is how IL-12 affects the function of macrophages especially independent of IFNγ. We hypothesize that IL-12 augments the ability of TIMs and TAMs to respond to inflammatory stimuli providing a window in which these stimuli are more likely to promote tumor destruction. Related to this hypothesis we wanted to investigate whether IL-12 changes signaling events associated with inflammatory signal transduction and determine if IFNγ required for all of the IL-12 induced changes in the response of TIMs and TAMs to inflammatory stimuli.

Herein, we use the highly aggressive and metastatic 4T1 mammary carcinoma murine model to look at the impact of IL-12 on the growth 4T1 tumors as well as isolate myeloid cells and study their in vitro response to LPS stimulation as well as tumor derived factors after exposure to exogenous, recombinant IL-12. We also examined the impact of pre-incubation of TIMs with IL-12 followed by LPS treatment on MAPK pathway signaling. Finally, we sought
to determine if IFNγ was compulsory for the increased inflammatory profile and increased MAPK phosphorylation in TIMs exposed to IL-12 followed by LPS.

This work provides insight into role of IL-12 in the context of IFNγ and the MAPK pathway, both of which have broad influence in immunology and cancer. This study offers a corollary to the classically identified mechanisms of IL-12 in innate immunity and cancer immunotherapy.

*Note: In this dissertation, TIMs refers to tumor infiltrating myeloid cells and TAMs refers to tumor associated myeloid cells. Previous works refer to TIMs and TAMs as tumor infiltrating macrophages and tumor associated macrophages. We designated TIMs as being directly from within the tumor while TAMs are from distal sites of a tumor bearing mice. Some studies do not make this distinction and use TAMs designation for any macrophage or myeloid cell from within the tumor as well as other tissue sites in a tumor bearing animal. We use the designation of myeloid cells because portions of the pooled CD11b+ cells used in experiments displayed markers for macrophages (F4/80+) or MDSCs (Gr1+).
CHAPTER II

IL.12 REDUCES THE IN VIVO GROWTH OF THE 4T1 MAMMARY CARCINOMA

Introduction

Breast cancer is the most common cancer diagnosis and the second most common cause of cancer deaths among women in the United States (181). The incidence of breast cancer rose from 1973-1990, but then began to decline, while mortality from breast cancer remained steady until recently as annual decreases in mortality have been observed (181). In spite of these statistics, it is well known that breast cancer prognosis is bleak and mortality increases when metastases are present, an indication of late-stage disease (182). The 5-year survival rate for stage IV breast cancer is only 15% (182). The poor clinical outcomes that accompany late-stage breast cancer provide an impetus to identify new and effective treatment strategies, especially ones that focus on disrupting immunosuppression and inhibiting metastatic processes associated with breast cancer.

In this chapter, we observed the growth of the 4T1 mammary carcinoma in syngeneic BALB/c mice and measured the impact of IL-12 microspheres on tumor progression. Fred Miller and colleagues first isolated the spontaneously
occurring mouse 4T1 mammary carcinoma (183). It reliably metastasizes to lung, liver, and brains when implanted (183).

As previously stated, initial cancer clinical trials in human using IL-12 were unsuccessful due in large part to systemic cytotoxicity and short half-life of free cytokine (184). In our in vivo experiments, we use recombinant IL-12 encapsulated in polylactic acid to facilitate slow, sustained release of IL-12, and to circumvent IL-12-associated cytotoxicity.

Prior studies in our laboratory have demonstrated that IL-12 has a profound impact on tumor associated macrophages in the Lewis lung carcinoma (LLC) mouse model (180). Therefore we sought to expand our understanding of IL-12’s impact on TIMs and TAMs in a breast cancer cell line. Because the 4T1 cell line is a syngeneic mice model with consistent metastasis, it is a good system to monitor the role of macrophages in both early stages of tumor initiation as well later metastatic events. Additionally, this model is an excellent platform to study agents such as IL-12 that modulate the immune system with regards to the tumor microenvironment.

There is current interest in developing therapeutic cancer vaccine as opposed to prophylactic vaccine such as those associated with preventing human papilloma virus (HPV), which subsequently prevents cervical carcinoma. One approach to developing therapeutic vaccines is the fusion of dendritic cells and tumor cells. Dendritic cells are innate immune cells that are the primary antigen-presenting cells (APCs) within the immune system. Fusion of dendritic cells and tumor cells allows for combination of antigen presentation machinery,
such as CD80, CD86, and MHC molecules, and tumor-associated, and tumor-specific antigen (185). This fusion optimizes tumor antigen presentation through both the class I and II MHC pathways to generate potent immune responses from adaptive immune cells, such as CD8$^+$ and CD4$^+$ T cells (185). Clinical trials using patient bone marrow-derived dendritic cells and autologous cancer cells have been successful in treating cancers (186-188). This aim looks to confirm the anti-tumor effects of IL-12 in the 4T1 breast cancer model, as well begin to understand if and how IL-12 may impact the efficacy of a dendritic cell fusion vaccine.

**Methods and Materials**

**Mice and cell lines**

Seven to eight week old female BALB/c mice were obtained from Jackson Laboratories. All animal care was in compliance with relevant protocols, guidelines, and regulations approved by the University of Louisville Institutional Animal Care and Use Committee. 4T1 cells were obtained from ATCC. Tumor cells were maintained through in vivo passage with limited intermittent in vitro culture.

**In vivo tumor injections**

BALB/c mice were injected subcutaneously near the left, rear mammary fat pad with 2x10$^4$ 4T1 cells. Tumors were measured at 3-day intervals with calipers at two bisecting diameters. An approximate volume was calculated using
the formula \((\text{Length} \times \text{Width}^2) \times 6/\pi\). Tumor weights were also measured upon harvest.

**IL-12 treatment**

Microspheres containing 250ng/mg of IL-12 or empty control microspheres were prepared by phase inversion nanoencapsulation and were supplied by Dr. Nejat Egilmez (University of Louisville School of Medicine). Microspheres were resuspended at 2mg/100\(\mu\)L in sterile phosphate buffered saline (PBS). At the indicated time point, mice received one intratumoral injection of 4mg of IL-12 or control microspheres.

**Lung harvest and staining**

The lungs of 4T1-bearing BALB/c mice were perfused with 30mL of Dulbecco’s PBS (Cellgro) supplemented with 2% fetal bovine serum (FBS) (Atlanta Biologicals) using a 30mL syringe and a 20G1/2 needle. Lungs were placed in Omnisette tissue cassettes (Fisher Scientific), and the cassettes were submerged in 10% formalin solution, neutral buffered (Sigma) for 24 hours. Lung containing cassettes were then added to 70% ethanol for further processing. Lungs were paraffin-embedded and sections were stained with Hematoxylin Eosin (Embedding and staining performed by the University of Louisville Special Procedures Laboratory).
Bone marrow derived dendritic cells (BMDCs) cell generation

The femurs and tibias of BALB/c mice were flushed with DPBS+2%FBS using a 1mL syringe and 26Ga needle. Cells were passed through a nylon filter (Bellco), and red blood cells were lysed using Gey’s solution. Cell were plated in 100mm dishes in RPMI 1640 (Hyclone) containing, 5% FBS, 1M HEPES (Sigma), 10mg/mL Gentamicin sulfate (Sigma), 20ng/mL GM-CSF (R&D Systems), and 1:100 Glutamax (Invitrogen) for 7 days at 37°C. Media was refreshed at days 2 and 5.

4T1 and BMDCs fusion and selection

4T1 tumor cells and BMDCs were mixed at a ratio of 1:5 in FBS-free RPMI 1640 containing 1M HEPES and 10mg/mL Gentamicin sulfate and polyethylene glycol (PEG) (Sigma Hybrimax) at 37°C. CD11c positive cells were selected using EasySep mouse CD11c+ selection kit and magnet (StemCell Technologies). Cells were plated for 1 week and loosely adherent fused cells collected for further use.

Flow cytometry

Cells were stained with 1ug/10^6 cells anti-CD32/CD16 Fc block (BD Pharmingen). Cells were then stained with 0.2ug/uL anti-CD326 APC (eBioscience), rat anti-IgG PE (BD Pharmigen) and rat anti-IgG2a APC
Acquisitions were performed using FACScaliber (BD Biosciences) and analysis was done using Flowjo (Tree Star).

**Statistics**

Significant differences between groups were determined using 2-tailed student’s T test.

**Results**

**In vivo administration of IL-12 microspheres diminishes 4T1 growth and lung metastasis**

We aimed to validate that microspheres loaded with recombinant IL-12 reduced in vivo tumor growth and metastasis in the 4T1 model of breast cancer. Our results demonstrate that a single intratumoral injection of IL-12 microspheres precipitated significant decreases in tumor growth. Injection of IL-12 microspheres at day 12 resulted in a significant decrease in tumor weights (Figure 1). The growth curve of tumors injected with IL-12 microspheres at day 5 was significantly altered with IL-12 treated tumors showing a decreased rate of expansion (Figure 2). We were also interested in validating the impact of IL-12 on 4T1 tumor metastasis. We observed that injection of IL-12 microspheres at day 9 had a substantial effect on tumor metastasis as lungs from 4T1-bearing animals had diminished metastasis when compared to controls (Figure 3).
Polyethylene glycol facilitates 4T1 and dendritic cell fusion

Additionally, we sought to fuse 4T1 cell and dendritic cells to investigate how IL-12 affects the efficacy of a fusion vaccine. Our long-term interests include investigating whether IL-12 induced changes in the macrophages contribute to enhancing the efficacy of a tumor dendritic cell vaccine. Initial attempts at 4T1 and dendritic cells fusion using polyethylene glycol (PEG) were successful. We were able to facilitate approximately 27% fusion of CD11c⁺ dendritic cells with 4T1 cells as indicated by cells double positive for the epithelial marker, CD326 and CD11c (Figure 4).

Discussion

Results in this aim demonstrate that in vivo treatment with IL-12 reduces both 4T1 growth and metastasis. Also, our data illustrates that exposing 4T1 and dendritic cells to PEG is a reasonable method to facilitate adequate fusion of these cells for eventual use as a tumor vaccine in mice. One of the drawbacks of IL-12 cytokine therapy is its transient effect when used as a singular treatment (189, 190). Subsequent studies to understand the mechanisms by which IL-12 exerts its effects on TIMs and TAMs will assist in prolonging and exploiting the positive effects of IL-12. It is important that the window of decreased immunosuppression initiated by IL-12 be used to introduce anti-cancer treatments such as therapeutic vaccines.

TIMS and TAMs are clear mediators of the profound immunosuppression within the tumor microenvironment. Studies directly linking role of TIMS and TAMs in cancer immunosuppression and obstacles to effective cancer
therapeutic vaccination are important to expanding the number effective immunotherapeutic options available. Studies have shown that depletion of TIMs and TAMs improved the effects of chemotherapeutic agents (191-193), but little data exist directly showing that changing the functional profile of myeloid cells within the tumor microenvironment is sufficient and/or necessary to positively impact the efficacy of chemotherapeutic agents or therapeutic vaccines. Work in this aim is a precursor for future studies that will investigate the potential roles of TIMs, TAMs, and IL-12 in establishing long-term cancer remission in response to therapeutic vaccination. Our future goals include ascertaining the in vivo duration of the change in myeloid cell function after exposure to IL-12. We would also like to examine whether IL-12 prompted myeloid cell conversion to a pro-inflammatory, anti-tumorigenic phenotype will improve the efficacy of a dendritic cell-based vaccine.
Figure 1. In vivo IL-12 administration decreases 4T1 tumor weights. BALB/c mice were challenged with $2 \times 10^4$ 4T1 cells. At day 12, mice were treated intratumorally with 4mg of control microspheres or microspheres containing 1000ng of rmIL-12. At day 26, tumors were resected and weighed. The average tumor weights were determined. n=7 for untreated group, n=8 for control group, n=7 for IL-12 group. *p<0.05 for untreated vs. IL-12 and control vs. IL-12. Error bars=SEM
Figure 2. In vivo IL-12 administration decreases the rate of 4T1 tumor growth. BALB/c mice were challenged with 2x10^4 4T1 cells. At day 5, mice were treated intratumorally with 4mg of microspheres containing 1000ng of rmIL-12 or 1000ng of blank microspheres as a control group. At set intervals, tumors were measured manually with calipers. Tumor volume was determined using the formula \( (L \times W^2) / 6 \pi \). Average tumor volume was calculated for each day. n=5 for control and IL-12 groups. \( p \text{ value}<0.05 \) for difference between groups at day 14, 17, and 19. Error bars=SEM.
Figure 3. In vivo IL-12 administration decreases 4T1 tumor metastasis.

BALB/c mice were challenged with $2 \times 10^4$ 4T1 cells. At day 9, mice were treated intratumorally with 4mg control microspheres or microspheres containing 1000mg of rmIL-12 or left untreated as a control group. Lungs were harvested at day 26. H&E staining was performed and lung metastasis visually observed. Results are representative images of 5 lungs per treatment group.
Figure 4. Polyethylene glycol (PEG) facilitates fusion of 4T1 tumor cells and bone marrow-derived dendritic cells. Bone marrow derived dendritic cells were fused with 4T1 tumor cells at a ratio of 1:5 (4T1:DC) using 50% PEG. Fused CD11c+ cells were positively selected. A.) Epithelial cell marker, CD326, but not dendritic cell marker, CD11c, is expressed on 4T1 cells. B.) CD11c, but not CD326 is expressed on dendritic cells. C.) CD11c+/CD326+ cells after fusion (upper right quadrant). Data is representative of 2 independent experiments.
CHAPTER III

IL.12 MODIFIES RESPONSE OF TUMOR ASSOCIATED AND TUMOR INFILTRATING MYELOID CELLS TO LPS VIA IFNγ INDEPENDENT AND DEPENDENT MECHANISMS

Introduction

The functional profile of macrophages and myeloid cells within tumors is an important determinant of the immune system’s ability to defend against the growth and spread of the tumor. A plethora of research has shown that macrophages play a multi-faceted role in the initiation and progression of cancers. Research has expanded our understanding of the link between inflammation and cancer. This new knowledge has evolved into a model that describes cancer-related inflammation through the intrinsic or extrinsic pathways (5, 194). The intrinsic pathway refers to changes in oncogenes and tumor suppressor genes that contribute to initiation of inflammation (195). Alterations in genes such as RET, RAS, and MYC lead to the elaboration of pro-inflammatory cytokines and growth factors (5). These cytokines and growth factors activate immune cells including macrophages within the vicinity, which perpetuates the inflammatory state and can eventually lead to tumor formation (5). Conversely, the extrinsic pathway describes external factors such as chronic infection or chemical and environmental exposures that promote inflammation (195). Chronic
infection increasing the risk for cervical cancer (194) or ultraviolet radiation associated inflammation ultimately progressing to skin cancer are examples of the extrinsic pathway of cancer related inflammation. The intrinsic and extrinsic pathways both lead to the activation of transcription factors such as NF-κB and STAT3 (194, 195). Macrophages and myeloid cells are critical in these formative steps in cancer related inflammation. Macrophages within this inflammatory milieu produce TNFα, IL-6, COX2, reactive nitrogen and oxygen species that have been shown to promote cancer formation in a number of ways. Reactive nitrogen and oxygen species promote cancer formation by inducing mutagenesis and genetic instability by disrupting the activity of p53 and enhancing the activity of DNA methyltransferases, which leads to errant transcription (196, 197). IL-6 and TNFα promote carcinogenesis by stimulating angiogenesis and cell proliferation (198).

Inflammation is tightly control process that integrates various signals that initially promote, but eventually down-regulate inflammatory mediators. In cancer-related inflammation, homeostatic controls are still present although they are likely deranged. Changes in the functional phenotype of the macrophage and myeloid derived cells are integral in the shift towards immunosuppression in the tumor microenvironment (199). After chronic inflammation has persisted and cancer has been established, signaling pathways in macrophages and myeloid cells are activated that aid in thwarting the activities of immunocompetent cells which allows for tumor cell evasion and escape. A recent report showed that the non-canonical Wnt protein, Wnt5a induces a tolerogenic phenotype in TAMs
through a negative feedback loop with NF-κB (200). This study also reported that Wnt5a leads to the accumulation of immunosuppressive macrophages in breast cancer patients (200). Others have shown that perturbations in NF-κB signaling such as the formation of non-canonical p50 homodimers may be pivotal in the macrophages immunosuppressive phenotype specifically via increased IL-10 secretion (201, 202).

It has been shown that TIMs from a mouse sarcoma produced significantly less pro-inflammatory IL-6, TNFα, and IL-1β and more anti-inflammatory cytokines IL-10 and TGFβ and IFN-inducible chemokines MCP1, CCL5, CXCL9, CXCL10, and CXCL16 than thioglycollate-elicited peritoneal macrophages in response to LPS (203). This observation is attributed to defective MyD88 activation of NF-κB, but intact stimulation of the MyD88-independent IRF3 and STAT1 signaling pathway in TAMs and TIMS (203). More recent studies suggest that TLR4 stimulation of TAMs results in enhanced ERK1/2 phosphorylation that induces IRAK-M expression, a known inhibitor of MyD88 (204).

The association between the presence of large tumor infiltrates of macrophages and myeloid cells and poor patient prognosis for several different cancers is widely recognized (100-102, 205). This observation initially prompted research that looked to eradicate macrophages and myeloid cells within the tumor environment to improve outcomes in cancer patients (191) (206). Additionally, reports showed that macrophages within the tumor microenvironment displayed the “M2” phenotype that is immunosuppressive and tumor supportive (199, 207). This information further promoted the view that
macrophage eradication was a plausible approach to alter the microenvironment and positively impact patient outcomes. Clodronate, a bisphosphonate initially used to decrease hypercalcemia in multiple myeloma patients, was adapted for use to systemically eliminate myeloid cells and macrophages. Although this approach yields reduced cancer immunosuppression, the use of clodronate is limited by its systemic toxicity (207). This approach also fails to appreciate the plastic nature of macrophages and myeloid cells and their pro-inflammatory potential. So rather than removing macrophages from the tumor microenvironment completely, efforts to understand how to exploit the manipulability of macrophages may be more suitable to cancer immunotherapeutic endeavors (4).

Our lab is interested in the impact of IL-12 on macrophages and myeloid cells within the tumor microenvironment. We propose that IL-12 facilitates the opening of an pro-immunological window in TIMS and TAMs which deems them more permissive to subsequent inflammatory stimuli. We have previously demonstrated that polarized TAMs and TIMs can be converted from immunosuppressive and tumor-supportive to pro-inflammatory and immunogenic in vivo and in vitro which may contribute to tumor eradication after exposure to IL-12 (180). In addition, work in this laboratory showed that in vivo introduction of IL-12 in tumor-bearing mice leads to rapid release of IL-15 for TIMs which is critical for cancer regression and is not dependent on the presence of NK cells (173). Others revealed that IL-12 enhances the ability of myeloid cells within the tumors to activate CD8+ T cell cytotoxicity (175). These reports demonstrate IL-
12 as a modulator of the functional phenotype of macrophages and myeloid cells in the tumor microenvironment.

The relationship between IL-12 and IFNγ is well established as previously discussed in the first chapter of this dissertation. IL-12 induction of IFNγ from NK cells, macrophages, and T cells is pivotal to adaptive immune responses (208). Studies have shown that IFNγ is very important in IL-12 induced changes in macrophages and IL-12’s overall anti-tumorigenic effects (175, 209). For example, a study showed that IL-12 induced IFNγ caused cell cycle arrest T-regulatory cells and enhanced tumor clearance (210). Less research has focused on any IFNγ independent IL-12 effects on macrophages and myeloid cells in the tumor microenvironment. In a series of experiments, Kerkar et al., showed that although IFNγ was required for IL-12 induced tumor regression, the host’s endogenous cells (immune cells included) ability to respond to IFNγ was only partially required (175). This report showed that when wild-type (WT) and IFNγ−/− CD8+ T cells transduced with melanoma antigen, p-mel TCR, and IL-12 were adoptively transferred into WT mice bearing B16 melanoma tumors, both the WT and IFNγ−/− p-mel/IL-12 T cells induced a similar degree of tumor regression. Conversely, in the IFNγ−/− mice bearing B16 melanoma, anti-tumor responses were significantly impaired with both WT and IFNγ−/− p-mel/IL-12 T cells. This showed that IL-12 induced IFNγ from endogenous cells in the tumor microenvironment was required to elicit anti-tumor response. Interestingly, when IFNgr−/− mice bearing B16 melanoma were injected with WT p-mel/IL-12 T cells there was only a partial reduction in IL-12 anti-tumor
effects. This suggests that although IFNγ production by endogenous cells is required for IL-12’s anti-tumor effects, the endogenous cell (including immune cells) response to IFNγ is not the sole determinant of IL-12 induced tumor regression. This data presents the possibility that IL-12 effects on endogenous cells including immune cells that are independent of IFNγ as well as IFNγ’s direct interaction with tumor cells may also be components of IL-12 anti-tumor mechanisms.

Although research has thoroughly investigated many components of the IL-12 signaling pathway such as JAK2, TYK2, and STAT4, not as much focus has been placed on other downstream mediators of IL-12 signaling, especially in the context of myeloid-derived cells. As discussed in the dissertation’s introduction, there are many signaling elements that play a role in determining how myeloid cells respond to inflammatory stimuli. These include NF-κβ, WNT, AKT, AMPK, and SOCS among others. One pathway that has been explored in the context of IL-12, but deserves more attention because of its broad influence over inflammatory pathways is the MAPK pathway. Most of the published data concerning IL-12 and the MAPK pathway addresses the pathway’s involvement in IL-12 synthesis in response inflammatory stimuli such as LPS. Very little work explores how IL-12 stimulation itself affects this pathway. It has been shown in the macrophage cell line, J774E, that IL-12 induces p38 phosphorylation, but not JNK or ERK1/2 (211). Zhang et al., demonstrated that in T cells, p38 activation is required for IL-12 induced IFNγ production (212). However, this requirement
has not been established in primary macrophages, macrophage cell lines, TIMS, TAMs, or MDSCs.

In this study, we hypothesize that IL-12 alters how TIMs and TAMs from the 4T1 mammary carcinoma respond to inflammatory stimuli. LPS is used as a model stimulus to demonstrate that IL-12 provides an immunological window that enhances the response of TIMS and TAMs to subsequent stimuli in a manner similar to vaccine adjuvants. Given the strong association between IL-12 and IFNγ immunological effects, we also focus on whether IFNγ is required for IL-12 induced changes in TIMs in response to LPS. Because several pathways contribute to the inflammatory profile of myeloid cells, we also sought to identify if IL-12 affects common signaling pathways associated with myeloid responses to LPS in order to begin to understand how this proposed immunological window is established. We evaluated if IL-12 impacted the status of several inflammatory intracellular signaling molecules including mitogen activated protein kinases (MAPKs) after LPS stimulation of 4T1 TIMs.

**Materials and Methods**

**Mice**

Wild type BALB/c and IFNγ-deficient BALB/c (C.12S7(B6)-ifng^{tm1Ts/J}) were obtained from Jackson Laboratories (Bar Harbor, Maine, USA). Animals were maintained in the University of Louisville’s Research Resources Facility. All animal care and experimental procedures were performed using protocols and
guidelines approved by the University of Louisville Institutional Animal Care and Use Committee.

**Tumor harvesting and cell isolation**

4T1 cells were obtained from ATCC (Manassas, Virginia, USA). Tumor cells were maintained through in vivo passage with limited intermittent in vitro culture. Seven-to-eight weeks-old BALB/c or IFNγ-deficient mice were injected with $2 \times 10^4$ 4T1 cells subcutaneously at the rear, left mammary fat pad. Tumors were harvested at approximately day 26. Tumors from WT and IFNγ-deficient mice were pooled for isolation of CD11b+ cells. Tumors were processed using Tumor Dissociation Kit and gentleMACS tissue dissociator (Miltenyi Biotec, Auburn, California, USA). CD11b+ cells were selected using anti-CD11b magnetic microbeads, LS columns and MACS magnet system (Miltenyi Biotec, Auburn, California, USA). For peritoneal cell harvest, peritoneal fluid was obtained using a 21G1 needle and DPBS containing 2% FBS. Red blood cells were lysed using a balanced salt solution. Cells were depleted of CD19+ and CD5+ followed by positive selection of CD11b+ using LS columns and the MACS magnet system. Cell isolation purity was confirmed by flow cytometry and exceeded 95% for both tumor and peritoneal cell isolation.

**Reagents**

LPS (Escherichia coli serotype O11:B4) was purchased from Sigma-Aldrich (St. Louis, Missouri, USA). Recombinant mouse IL-12 was obtained from
R&D Systems (Minneapolis, Minnesota, USA). Western blot detection of protein used the following primary antibodies: anti-phospho-p38 (Thr180/Tyr182), anti-p38, anti-phospho-JNK/SAPK (Thr183/Tyr185), anti-JNK/SAPK, anti-phospho-p44/42 (Thr202/Tyr204), anti-p44/42 (Cell signaling Technology, Beverly Massachusetts, USA) and anti-B-actin (Sigma-Aldrich, St. Louis, Missouri, USA).

**In vitro treatment protocol**

For **protein**, CD11b+ cells were treated with 100ng/mL LPS, 100ng/mL IL-12, or IL-12 + LPS at the same concentrations for 18 hours in RPMI with 5% fetal bovine serum. For IL-12 pretreatment, CD11b+ cells were treated with 100ng/mL IL-12 for 2 hours followed by 100ng/mL LPS for 18 hours. For **mRNA**, CD11b+ cells were treated with 100ng/mL LPS, 100ng/mL IL-12, or IL-12 + LPS at the same concentrations for 2 hours in RPMI with 5% fetal bovine serum. For IL-12 pretreatment, CD11b+ cells were treated with 100ng/mL IL-12 for 2 hours followed by 100ng/mL LPS for 2 hours. All incubations were done at 37˚C.

**Cytokine measurement**

Tumor infiltrating and peritoneal CD11b+ cells were harvested as described above. Following stimulation, TNFα, IL-6, and IL-10 protein levels were measured using the Cytometric Bead Array Mouse Inflammation Kit (BD Biosciences, San Jose, California, USA) according the manufacturer’s instructions. Analysis was performed using FACSCalibur cytometer and FCAP array software (BD Biosciences, San Jose, California, USA).
Real-time PCR analysis

MACs One-step cDNA Kit (Miltenyi Biotech, Auburn, California, USA) was
used for RNA and cDNA synthesis. cDNAs were amplified using a 20μL reaction
containing SYBR green (New England Biolabs, Ipswich, Massachusetts, USA).
QuantiTECT primers assays for mouse TNFα, IL-6, IL-10, and B-actin (Qiagen,
Hilden, Germany) were used and PCR reactions were performed and captured
by DNA Opticon 2 (MJ Research/Bio-rad, Hercules, California, USA). The
relative expression software tool (REST 2009) was used to calculate relative fold
changes among treatment groups.

Western blot analysis

Cells were lysed in buffer containing 125mM Tris (pH 6.8), 2% SDS, 20%
glycerol, 200μM PMSF, protease inhibitor mixture (Promega, Madison,
Wisconsin, USA), phosphatase inhibitor mixture (ThermoFisher Scientific,
Rockford, Illinois, USA). Total protein content for each sample was assessed by
BCA protein assay (ThermoScientific, Rockford, Illinois, USA). Equal amounts of
protein were separated on 10% Criterion gels (Bio-rad, Hercules, California,
USA), by SDS-PAGE. Proteins were transferred to nitrocellulose membranes
using Trans-blot Turbo Transfer system (Bio-rad, Hercules, California, USA).
Antibody-bound proteins were detected using an ECL western blot system (GE
Healthcare, Pittsburgh, Pennsylvania, USA), and the membranes were exposed
to UltraCruz 5x7 autoradiography film (Santa Cruz Biotechnology, Santa Cruz, California, USA)

**Flow cytometry**

Single cell suspensions of WT and IFNγ-deficient CD11b+ tumor infiltrating cells were stained with fluorescently conjugated antibodies against murine CD11b, F4/80, pan Gr1, Ly6G, Ly6C (all but anti-F4/80 obtained from BD Biosciences, San Jose, California, USA; anti-F4/80 obtained from Affymetrix/eBiosciences, San Diego, California, USA) for 30 minutes at 4°C, washed and analyzed using FACSCalibur flow cytometer and FlowJo software (Tree Star, Ashland, Oregon, USA).

**4T1 supernatant production**

Primary 4T1 cells were harvested directly from tumor bearing animals by mashing the tumor mass through a 100 micron filter cup and then passage over a 70 micron nylon filter were plated in 10 cm dishes overnight at 10^7 cells/dish in normal RPMI 1640 with 5% FBS. After overnight culture, tissue debris and stroma were removed from the plates by pipetting away the supernatant and replacing with fresh media. The cell cultures were maintained at 37°C and 5%CO₂ and allowed to grow to 70% confluency. At this point, cells were split and transferred for expansion. Once desired growth was achieved, supernatants were collected from the flasks and centrifuged at 450g for 5 minutes. The pellet was discarded and the clarified supernatants were then centrifuged a 2000g for
10 minutes. The supernatant was then applied to a Vivacell 100kd MWCO concentrator (Sartorius) and centrifuged in multiple 10-15 minute spins until the volume of the retentate was reduced to a final volume of approximately 3-4 ml. The retentate was aliquotted into 500 μl amounts and stored at -20°C until use. Concentrated retentate was tested for endotoxin by LAL (Limulus Amebocyte Lystate) Test (Lonza, QCL-1000 assay). The concentrate was also screened for mycoplasma using MycoAlert Assay (Lonza). Protein concentration was determined by BCA (Pierce). Optimal stimulation dose was determined by dose response curve and typically was 100-300 ng/ml. Verified concentrate was then used to stimulate macrophages in in-vitro assays.

**Statistical analysis**

Statistical significance was calculated using unpaired student’s t test when comparing two groups. One-way ANOVA was used for statistical difference when analyzing greater than two groups. p<0.05 was considered to be statistically significant.

**Results**

*IL-12 impacts the response of 4T1 tumor associated myeloid cells to LPS*

It has been reported that TIMs and TAMs from mice bearing established tumors are polarized towards a functional phenotype that promotes immunosuppression and immune evasion as well as metastasis (213, 214). Additionally, TIMs and TAMs have been implicated in decreasing the efficacy of chemotherapeutic agents (215). TIMs and TAMs express high levels of anti-
inflammatory cytokines, IL-10 and TGFβ, and low levels of pro-inflammatory cytokines such as TNFα, IL-12, IFNγ, and IL-6 (213) Introduction of a priming molecule that would allow for a more robust TIM and TAM response to a subsequent pro-inflammatory stimulus could help overcome this profound immunosuppression and even improve the effects of chemotherapeutic agents.

To test the hypothesis that IL-12 acts a primer and impacts the ability of TIMs and TAMs to respond to an inflammatory stimulus, CD11b+ cells were isolated from the peritoneum of 4T1 bearing BALB/c mice and from non-tumor bearing control BALB/c mice. These TAMs and control peritoneal myeloid cells were treated with IL-12 alone, LPS alone, IL-12 + LPS simultaneously for 18 hours, or pretreated with IL-12 for 2 hours followed by LPS for 18 hours. This treatment protocol allowed for examination of whether IL-12 immediately changed TAMs response to LPS or if there was a temporal effect seen with IL-12 pretreatment followed by LPS treatment.

We measured the levels of pro-inflammatory cytokines, TNFα and IL-6, and the anti-inflammatory cytokine, IL-10 using a cytometric bead array (CBA). We then compared our treatment groups in both 4T1 peritoneal TAMs (5A, D, G) and control peritoneal myeloid cells (Fig. 5B, E, and H). We also compared each treatment group between 4T1 peritoneal TAMs and control peritoneal myeloid cells side-by-side (Fig. 5C, F, and I).

We found that IL-12 alone did not elicit any TNFα from either control peritoneal myeloid cells or 4T1 peritoneal TAMs (Fig. 5A and B). LPS alone significantly increased the amount of TNFα secreted by control peritoneal
myeloid cells and 4T1 peritoneal TAMs (Fig. 5A and B). When directly comparing the amount of TNFα secreted by control peritoneal myeloid cells and 4T1 peritoneal TAMs in response to LPS, we show that there was an overall decrease in TNFα produced by 4T1 peritoneal TAMs, but this decrease was not significant (Fig. 5C). When peritoneal myeloid cells were treated with IL-12+LPS simultaneously, we again saw an increase in TNFα production as compared to untreated and IL-12 treated groups in both control peritoneal myeloid cells and 4T1 peritoneal TAMs (Fig. 5A and B). When we directly compared, IL-12+LPS treatment between control peritoneal macrophages and 4T1 peritoneal TAMs, there was no difference in TNFα cytokine production (Fig. 5C). When control peritoneal myeloid cells and 4T1 peritoneal TAMs were pretreated with IL-12 followed by LPS, we observed increased TNFα secretion over untreated and IL-12 treated control peritoneal myeloid cells and 4T1 peritoneal TAMs. There was no significant difference in the amount of TNFα produced after IL-12 pretreatment followed by LPS when we directly compared peritoneal myeloid cells from control and 4T1-bearing mice (Fig. 5C). There was a significant decrease in the amount of TNFα produced in response to IL-12+LPS when compared to LPS alone and IL-12 pretreatment followed by LPS in control peritoneal myeloid cells (Fig. 5A). Although not statistically significant, a similar trend was noticed in 4T1 peritoneal TAMs. Overall, pretreatment of control peritoneal myeloid TAMs and 4T1 peritoneal TAMs did not result in a significant increase in the amount of TNFα secreted in response to LPS. Simultaneously treatment of control myeloid cells and 4T1 peritoneal TAMs caused a decrease in the amount of TNFα produced
when compared to cells treated with LPS alone or IL-12 pretreatment followed by LPS.

We noted that when both control peritoneal myeloid cells and 4T1 peritoneal myeloid cells were exposed to LPS, IL-12+LPS, and IL-12 pretreatment followed by LPS, both cell types produced significantly higher amounts of IL-6 as compared to untreated and IL-12 treated cells (Fig. 5D and E). We also observed, although not statistically significant that pretreatment of peritoneal myeloid cells from both control and 4T1-bearing mice with IL-12 followed by LPS enhanced the amount of IL-6 produced as compared to LPS or IL-12+LPS treatment (Fig. 5A and B). When compared directly, control peritoneal myeloid cells and 4T1 peritoneal TAMs secreted similar amounts of IL-6 in response to each treatment (Fig. 5F).

Control peritoneal myeloid cells and 4T1 peritoneal TAMs secreted similar amounts of IL-10 in response to LPS alone and IL-12+LPS which was a significant increase over cells that were untreated or treated with IL-12 alone (Fig. 5G and H). Interestingly, both control peritoneal myeloid cells and 4T1 peritoneal TAMs secreted significantly less IL-10 after being pretreated with IL-12 followed by LPS as compared to treatment with LPS alone or IL-12+LPS (Fig. 5G and H). When we compared each treatment group directly between control peritoneal myeloid cells and 4T1 peritoneal TAMs, there was a significant difference in the amount IL-10 produced in response to IL-12+LPS (Fig. 5I). Overall, pretreatment with IL-12 followed by LPS treatment of control peritoneal
myeloid cells and 4T1 peritoneal TAMs resulted in a significant decrease in the amount of IL-10 produced as compared to LPS alone and IL-12+LPS.

4T1 peritoneal TAMs produce less IFNγ than control peritoneal myeloid cells in the presence of IL-12.

Because of the strong association between the immune effects of IL-12 and IFNγ, we observed the amount of IFNγ produced by control peritoneal myeloid cells and 4T1 peritoneal TAMs. We wanted to determine if there was a difference in the amount of IFNγ produced by control peritoneal myeloid cells and 4T1 peritoneal myeloid cells. When we looked at mRNA levels of IFNγ, we noted that all treatments of both control peritoneal myeloid cells significantly affected the level of IFNγ mRNA expression (Fig 6A and B). First, we compared the mRNA levels when each treatment group for control peritoneal myeloid cells and 4T1 peritoneal TAMs was normalized to untreated cells of the same type (Fig. 6A). We found that there was significantly more IFNγ mRNA expressed in control peritoneal myeloid exposed to IL-12, IL-12+LPS, and IL-12 pretreatment followed by LPS as compared to 4T1 peritoneal TAMs. Secondly, we compared the IFNγ mRNA levels between control peritoneal myeloid cells and 4T1 peritoneal TAMs when all treatments for both cell types were normalized to untreated control peritoneal myeloid cells (Fig. 6B). We found that 4T1 peritoneal TAMs expressed significantly less IFNγ mRNA as compared to control peritoneal myeloid cells (Fig. 6B), and this observation was unchanged with the addition of IL-12 alone or LPS alone. The addition of IL-12 simultaneously with LPS or as pretreatment
before LPS significantly increased the amount of IFNγ mRNA in both control peritoneal myeloid cells and 4T1 peritoneal TAMs (Fig 6A). There was still, however, a large difference in the amount of IFNγ mRNA between control peritoneal myeloid cells and 4T1 peritoneal TAMs in response to IL-12+LPS and IL-12 pretreatment followed by LPS when untreated control peritoneal myeloid cells were used as the reference (Fig. 6B).

We also compared IFNγ cytokine produced in response to each treatment in control peritoneal myeloid cells and 4T1 peritoneal myeloid cells. We found that IL-12 alone and LPS alone resulted in a small increase in IFNγ secretion in both control peritoneal myeloid cells and 4T1 peritoneal TAMs, and that there was no difference between the two groups with either treatment. When myeloid cells from the peritoneum of control and 4T1-bearing mice were exposed to IL-12+LPS and IL-12 pretreatment followed by LPS, they both secreted significantly more IFNγ as compared to treatment with IL-12 or LPS (Fig. 6C). 4T1 peritoneal TAMs secreted significantly less IFNγ in response to IL-12+LPS and IL-12 pretreatment followed by LPS (Fig. 6C). From these data, we show that 4T1 peritoneal TAMs express less IFNγ mRNA and cytokine levels at baseline. The addition of IL-12+LPS and IL-12 pretreatment followed by LPS increases the amount of IFNγ produced in 4T1 peritoneal TAMs, but not to the same extent as in control peritoneal myeloid cells.

Absence of IFNγ affects the response of 4T1 TAMs to IL-12 and LPS
Next, we evaluated whether the absence of IFNγ would affect the production of TNFα, IL-6, and IL-10 by 4T1 peritoneal TAMs. 4T1 TAMs were isolated from the peritoneums of IFNγ-deficient mice and treated as mentioned above. We compared the mRNA and proteins levels of each cytokine from wild-type and IFNγ-deficient 4T1 peritoneal TAMs. We found 4T1 peritoneal TAMs from IFNγ-deficient mice secreted significantly less TNFα in response to LPS, IL-12+LPS, and IL-12 pretreatment followed by LPS as compared to wild-type 4T1 peritoneal TAMs (Fig 7A). The level of TNFα mRNA expression was consistently lower in IFNγ-deficient 4T1 peritoneal TAMs than in wild-type 4T1 peritoneal TAMs for treatment with LPS, IL-12+LPS, and IL-12 pretreatment followed by LPS although this difference was only statistically significant for IL-12 pretreatment followed by LPS (Fig. 7D).

Wild-type 4T1 peritoneal TAMs secreted slightly more IL-6 in response to LPS, IL-12+LPS, and IL-12 pretreatment followed by LPS than IFNγ-deficient 4T1 peritoneal TAMs, but there was no significant difference in the values (Fig 7B). When comparing IL-6 mRNA expression between the two groups, we observed that IFNγ-deficient 4T1 peritoneal TAMs had increased IL-6 mRNA after stimulation with LPS, IL-12+LPS, and IL-12 pretreatment followed by LPS (Fig. 7E).

Finally, our data shows that IFNγ-deficient 4T1 peritoneal TAMs produced significantly more IL-10 in response to LPS, IL-12+LPS, and IL-12 pretreatment followed by LPS as compared to wild-type 4T1 peritoneal TAMs (Fig 7C). Additionally, IL-10 mRNA levels were higher in IFNγ-deficient 4T1 peritoneal
TAMs relative to wild-type 4T1 peritoneal TAMs in response to LPS, IL-12+LPS, and IL-12 pretreatment followed by LPS (Fig. 7F).

It should also be noted that if the cytokines data for IFNγ-deficient 4T1 peritoneal TAMs is observed in isolation, pretreatment with IL-12 results increased TNFα and IL-6 and decreased IL-10 in response to LPS as compared to LPS alone and IL-12 + LPS treatment (Figs. 7A-C). The pattern was retained in IFNγ-deficient 4T1 peritoneal TAMs mRNA expression data for IL-6 (Fig. 7E) and IL-10 (Fig. 7F), but lost for TNFα (Fig. 7D).

Overall, this data demonstrates that IFNγ is important for IL-12 induced changes in TNFα production in response to LPS in 4T1 peritoneal TAMs. Conversely, our data suggests that IFNγ is not necessary for the modification of IL-6 gene and cytokine expression seen with IL-12 pretreatment followed by LPS, and may even negatively influence IL-6 production in 4T1 peritoneal TAMs. We also show that IFNγ is not required for decreased IL-10 production seen with IL-12 pretreatment followed by LPS in 4T1 peritoneal TAMs, even though IFNγ-deficient 4T1 peritoneal TAMs produced more IL-10.

*IL-12 modulates the response of 4T1 tumor-infiltrating myeloid cells to LPS.*

We also tested whether IL-12 impacts the ability of myeloid cells to respond to an inflammatory stimulus by using CD11b+ cells isolated from 4T1 tumors. These 4T1 CD11b+ tumor infiltrating myeloid cells (4T1 TIMs) were then exposed to the following treatments in vitro: IL-12 alone, LPS alone, IL-12 plus LPS simultaneously, or IL-12 pretreatment followed by LPS. Cytokine production was
measured using a cytometric bead array (CBA). We observed that IL-12 alone does not induce significant production of TNFα, IL-6, or IL-10 (Fig. 8A-F) or their respective mRNAs (Fig. 8A, C, E). Conversely, LPS alone and IL-12 plus LPS given simultaneously, resulted in significant production of TNFα, IL-6, and IL-10 (Fig. 8A, C, E) and their respective mRNAs (Fig. 8A, C, E). When 4T1 TIMs were pretreated with IL-12 and then exposed to LPS, we observed a significant increase in the protein and mRNA levels of TNFα and IL-6 as compared to treatment with LPS alone or IL-12 plus LPS (Fig. 8A-D). On the other hand, pretreatment with IL-12 followed by LPS treatment of 4T1 TIMs resulted in a significant decrease in the amount of IL-10 (Fig. 8E-F).

We also assessed if IL-12 increased the amount of IFNγ produced by 4T1 TIMs in response to LPS. We show that IL-12 induced a significant amount of IFNγ mRNA and cytokine from 4T1 TIMS as compared untreated 4T1 TIMs (Fig. 9A and B). 4T1 TIMS produced significantly more IFNγ when treated with IL-12 +LPS or pretreated with IL-12 followed by LPS than when treated with LPS alone (Fig. 9A and B).

*IL-12 increases pro-inflammatory cytokine produced by 4T1 TIMs in response to 4T1 tumor derived products*

We have demonstrated that IL-12 enhances the production of pro-inflammatory cytokines, IL-6 in 4T1 peritoneal TAMs, TNFα and IL-6 in 4T1 TIMs and decreases the secretion of IL-10 by 4T1 peritoneal TAMs and 4T1 TIMs in response to LPS. We used LPS as a model stimulus to demonstrate that IL-12
augments how TIMs and TAMs respond to inflammatory stimuli. It has been proposed that the tumor secrete products with pro-inflammatory and anti-tumorigenic potential but because of chronic exposure to these signals, immunosuppression is induced leading to continued immune evasion, tumor growth, and metastasis (216). We investigated whether IL-12 could also augment the response of 4T1 TIMs to tumor derived products. We concentrated supernatants from ex vivo harvested 4T1 tumors, and treated 4T1 TIMs with IL-12 or 4T1 supernatant alone, IL-12+4T1 supernatant, IL-12 pretreatment followed by 4T1 supernatant. We found that treatment with 4T1 supernatant led to a small but significant amount of TNFα and IL-6 from 4T1 TIMs (Figs. 10A and 10B). We also noted the 4T1 supernatant alone caused a tremendous increase in the amount IL-10 secreted from 4T1 TIMs (Fig. 10C). A major finding was that IL-12 pretreatment led to a significant enhancement of the amount of pro-inflammatory cytokines, TNFα and IL-6, produced in response to 4T1 supernatants (Figs. 10A and B). Interestingly, treating 4T1 TIMs with IL-12+4T1 supernatants simultaneously or pretreating with IL-12 followed by 4T1 supernatant did affect the amount of IL-10 produced (Fig. 10C). Overall, this data shows that IL-12 changes the response of 4T1 TIMs to 4T1 supernatant from predominantly IL-10 to a phenotype that includes proinflammatory cytokines, TNFα and IL-10 in addition to IL-10.

*IL-12 modulates the response of IFNγ-deficient, 4T1 tumor-infiltrating myeloid cells to LPS.*
IFNγ is well known activator of myeloid cells and IL-12 has been shown to induce IFNγ production by myeloid cells (217). Additionally, it has been shown that much of IL-12’s anti-tumor influence can be attributed to IFNγ (175, 210, 218). Conversely, there are studies that demonstrate that IL-12 has immune properties that are independent of IFNγ (219, 220). Therefore, we sought to determine if IFNγ was important for our observed finding that pretreatment of 4T1 TIMs with IL-12 increased the amount of pro-inflammatory cytokines, TNFα and IL-6 and decreased anti-inflammatory cytokine, IL-10 produced in response to LPS. To approach this question, 4T1 TIMs were isolated from IFNγ-deficient mice and treated in the same manner as their wild-type counterparts.

To determine if there were differences in the myeloid cell population of 4T1 tumors harvested from wild-type and IFNγ-deficient mice, CD11b+ cells from the tumors of both animal types were analyzed using flow cytometry. The level of expression of myeloid cell markers Gr1, F4/80, Ly6C, and Ly6G were insignificant between wild-type and IFNγ-deficient 4T1 TIMs (Fig 11).

When IFNγ-deficient 4T1 TIMs were stimulated, we found that IL-12 alone stimulated minimal TNFα, IL-6, and IL-10 (Fig. 12A-F). IFNγ-deficient 4T1 TIMs produced similar amounts of TNFα, IL-6, and IL-10 in response to LPS and IL-12 plus LPS (Fig. 12A, C, E). IL-12 pretreatment followed by LPS exposure caused a significant increase in TNFα and IL-6 cytokine production and a reciprocal decrease in IL-10 cytokine production by IFNγ-deficient 4T1 TIMs (Fig. 12A, C, E). However, no differences in gene expression were observed between LPS alone, IL-12 plus LPS, and IL-12 pretreatment followed by LPS in IFNγ-deficient
Overall, the absence of IFNγ in 4T1 TIMs did not significantly affect the increase in TNFα and IL-6 cytokine levels and reciprocal decrease in IL-10 seen with IL-12 pretreatment followed by exposure to LPS. Conversely, there was no increase in TNFα and IL-6 gene expression and no decreased IL-10 gene expression seen in IFNγ-deficient 4T1 TIMs pretreated with IL-12 followed by LPS. This data suggest that the priming effects of IL-12 works through various mechanisms at both the transcriptional and translational levels, and may have an IFNγ independent component.

*IFNγ partially impacts the response of 4T1 TIMs to IL-12 and LPS*

The data previously described shows cytokine and gene expression when comparing our various treatment groups in either wild-type 4T1 TIMs exclusively or IFNγ-deficient 4T1 TIMs exclusively. This data shows that even in the absence of IFNγ, pretreatment of 4T1 TIMs with IL-12 increases the amount of TNFα, IL-6 cytokine produced, and decreases the amount of IL-10 cytokine produced in response to LPS. Conversely, we demonstrate that increased TNFα and IL-6 gene expression and decreased IL-10 gene expression induced by IL-12 pretreatment followed by LPS is no longer present in IFNγ-deficient 4T1 TIMs. These observations prompted us to directly compare each treatment group between wild-type and IFNγ-deficient 4T1 TIMs to more clearly understand the influence of IFNγ. To directly compare each treatment between WT and IFNγ-deficient 4T1 TIMs, we reanalyzed our gene data with both wild-type and IFNγ-deficient 4T1 TIMs mRNA levels for each gene normalized to wild-type untreated
4T1 TIMs. At baseline, we found that there was no difference in TNFα gene expression between untreated wild-type and IFNγ-deficient 4T1 TIMs (Fig. 13A). There was, however, a small but significant difference in IL-6 gene expression between untreated wild-type and IFNγ-deficient 4T1 TIMs that was not affected by IL-12 treatment (Fig 13B). Additionally, there was significantly less IL-10 gene expression in untreated IFNγ-deficient 4T1 TIMs as compared to wild-type 4T1 TIMs (Fig. 13C). This difference in IL-10 mRNA levels between wild-type and IFNγ-deficient 4T1 TIMs was decreased with IL-12 treatment (Fig 13C). We found that TNFα gene expression was significantly lower in the IFNγ-deficient 4T1 TIMs as compared to wild-type 4T1 TIMs in response to LPS alone, IL-12 plus LPS, and IL-12 pretreatment followed by LPS (Fig. 13A). There was no significant difference in IL-6 gene expression between wild-type and IFNγ-deficient 4T1 TIMs treated with LPS, IL-12 plus LPS, and IL-12 pretreatment followed by LPS (Fig. 13B). For IL-10 gene expression, there was no difference in the IL-10 gene response of wild-type and IFNγ-deficient TIMs to IL-12 pretreatment followed by LPS, but there was a significant difference in IL-10 gene expression between wild-type and IFNγ-deficient 4T1 TIMs in response to IL-12 plus LPS (Fig. 13C). Although the difference in IL-10 gene expression in response to LPS was not significant between wild-type and IFNγ-deficient TIMs, there was noticeable variance between the two TIM populations (Fig. 13C).

To directly compare cytokine levels of wild-type and IFNγ-deficient TIMs in response to our treatments, we combined cytokine data from figures 1 and 2 in table form. When directly comparing cytokine production between wild-type and
IFNγ-deficient 4T1 TIMs for each treatment, we observed lower amounts of TNFα in IFNγ-deficient 4T1 TIMs for all treatment groups although there was only a significant decrease in TNFα for the IL-12 pretreatment followed by LPS group (Table I). There was no significant difference in IL-6 cytokine production between wild-type and IFNγ-deficient 4T1 TIMs for any of the treatment groups (Table I). Finally, although there was an overall decrease in IL-10 cytokine production in IFNγ-deficient 4T1 TIMs for all treatment groups, there was only a significant difference in the IL-12 treatment group when comparing the two populations (Table I).

In summary, by directly comparing gene expression and cytokine production in wild-type and IFNγ 4T1 TIMs, we show that IFNγ is necessary for the increase in TNFα after IL-12 pretreatment followed by LPS. Conversely, our data demonstrate that the effect of IL-12 pretreatment on the response of 4T1 TIMs to LPS in terms of IL-6 gene expression and cytokine production is independent of IFNγ. Finally, we show that IL-12 decreases the amount of IL-10 cytokine produced, but not IL-10 gene expression by 4T1 TIMs in the absence of IFNγ.

*IL-12 alters the 4T1 TIMs activation of MAPK pathway in response to LPS*

Our data thus far has shown that IL-12 affects the response of TIMs to LPS in a manner that is partially independent of IFNγ. We observed the levels of TNFα, IL-6, and IL-10 in response to IL-12 and LPS. There are many pathways involved in the production of these cytokines in myeloid cells in response to inflammatory
stimuli such as LPS including signal transducer and activator of transcription (STATs), NF-κβ, AP-1, and AKT (12). The MAPK signaling cascade is well-studied and has been shown to be important in the signaling pathway of TNFα, IL-6, IL-10, and LPS (81, 87, 90). These same signaling pathways also trigger the resolution of inflammation through feedback inhibition loops that lead to activation of suppressor of cytokine signaling (SOCS) proteins and IL-10 production (12). IL-12 signaling is propagated via JAK2 and STAT4 (156, 157). It has also been reported that IL-12 activates the MAPK pathway (161). We hypothesized that IL-12 may affect the activation status of one or more of these pathways to augment the inflammatory profile of 4T1 TIMs in response to LPS. Additionally, given the evidence that the MAPK pathway is involved in both IL-12 and LPS signaling, we postulated that the MAPK pathway may be affected in 4T1 TIMs exposed to IL-12 followed by LPS.

First, we investigated the impact of IL-12 on several inflammatory signaling components. We found that IL-12 did not affect the phosphorylation status of NF-κβ p65 in 4T1 TIMs (Fig. 14A) In addition to this finding, we found no change in total amount of NF-κβ p50 in 4T1 TIMs (Fig 14A). We also looked at the influence of IL-12 on 4T1 TIM expression of cyclooxygenase-2 (COX-2), which has been shown to be important in cancer-related inflammation (89). We observed no change in 4T1 TIM expression of COX-2 in response to IL-12. We also noted that IL-12 did not affect the phosphorylation of GSK3β and AKT in 4T1 TIMs (data not shown).
Next, we evaluated the effect of IL-12 on the MAPK pathway. For these experiments, we also gauged how pretreatment with IL-12 affected the MAPK pathway after exposure to LPS. When wild-type 4T1 TIMs were exposed to LPS alone, there was phosphorylation of MAPK proteins p38, JNK, and p44/42 that diminished over time (Fig. 14B). Conversely, IL-12 alone elicited a very low level of phosphorylation of the MAPK protein p38, but no p44/42 or JNK phosphorylation at the observed time points (Fig. 14B). When TIMs were pretreated with IL-12 for 2 hours and then exposed to LPS, there was significant enhancement of phosphorylation of p38, JNK, and p44/42 as compared to IL-12 or LPS alone (Fig. 14B).

Finally, we examined MAPK protein phosphorylation in IFNγ-deficient 4T1 TIMs in response to IL-12 and LPS. We wanted to assess if the enhancement of MAPK signaling seen with IL-12 pretreatment of 4T1 TIMs was at all IFNγ independent. We observed that treatment of IFNγ-deficient 4T1 TIMs with LPS alone resulted in a similar pattern of phosphorylation of p38, JNK, and p44/42 as seen in wild-type 4T1 TIMs (Fig. 14C). IL-12 caused some phosphorylation of p38, as well as p44/42 MAPK proteins in IFNγ-deficient 4T1 TIMs (Fig. 14C). Finally, pretreatment of IFNγ-deficient 4T1 TIMs with IL-12 did not cause a notable increase in p38 and JNK phosphorylation, as seen in wild-type 4T1 TIMs (Fig. 14C). However, pretreatment of IL-12 followed LPS treatment did cause increased phosphorylation of p44/42 at an earlier time point (15 minutes) when compared to LPS alone (Fig. 14C).
Overall we demonstrate that IL-12 does not impact activation or expression of NF-κβ in 4T1 TIMs. Additionally, the pro-inflammatory enzyme, COX-2 was not affected by IL-12 treatment of 4T1 TIMs. We do show that IL-12 enhances the phosphorylation of MAPK proteins after LPS stimulation in 4T1 TIMs. Our data suggests that the enhanced phosphorylation of p38 and JNK is dependent on IFNγ as this enhancement was lost when IFNγ-deficient 4T1 TIMs were pretreated with IL-12 followed by LPS. We did, however, identify that in the absence of IFNγ, IL-12 enhances the phosphorylation of p44/42 in response to LPS in 4T1 TIMs at an earlier time point.
Discussion

Myeloid-derived cells are remarkably dynamic and display an array of functions and activities. The dynamic nature of myeloid cells is especially evident in cancer. Myeloid cells in cancer have the ability to initiate and promote tumor growth, as well as suppress their own and other cell types’ capability to thwart cancer progression which provides an opportunity for targeting these cells in anti-cancer therapies. Many have sought ways to exploit the functional plasticity of myeloid cells in cancer in order to aid in eradicating established tumors. Recent approaches that have been shown to modulate the functional phenotype of TAMs and TIMs and improve anti-tumor immunity include engagement and activation of TAMs and TIMs via TLR4 (221), local low-dose irradiation (222), and administration of biological nanoparticles combined with chemotherapeutic agents (223). IL-12 can activate both the innate and adaptive arms of the immune system and much effort has been used to investigate its effectiveness as an anti-tumor agent and therapeutic vaccine adjuvant. In this study, we explore how IL-12 impacts the tumor associated and tumor infiltrating myeloid cell’s response to the inflammatory stimulus, LPS, as well as tumor derived products. We use LPS solely as a model inflammatory stimulus to demonstrate that IL-12 produces an immunological window in which TAMs and TIMs are more able to respond to subsequent pro-inflammatory and anti-tumor signals. Interestingly, it has been shown that ex-vivo treatment with LPS of tumor-associated macrophages from the ascites and blood of ovarian cancer patients leads to enhances activation and tumor cell cytoxicity of NK cells (221).
Our data show that IL-12 enhances the amount of TNFα and IL-6 and decreases the amount of IL-10 4T1 TIMs secrete in response to LPS (Fig. 8). This result aligns with data using TIMs from the 3LLC lung cancer (180). An important distinction in this study include showing that exposure of TIMs for only 2 hours versus overnight pre-culture with IL-12 conferred a similar cytokine response to LPS in vitro. Additionally, treating TIMs with IL-12 and LPS simultaneously did not impact the cytokine and mRNA levels to the degree that pre-treatment with IL-12 followed by LPS treatment. This observation suggests that although IL-12 does not induce the production of cytokines and mRNA measured in TIMs, it does initiate a temporal and internal change in myeloid cells that alters its response to subsequent stimuli.

In 4T1 peritoneal TAMs, adding IL-12 simultaneously with LPS led to a slight decrease in TNFα secretion (Fig. 5B). This difference was no longer present when 4T1 peritoneal TAMs were pretreated with IL-12 followed by LPS (Fig 5B). We saw an increase in IL-6 cytokines levels when 4T1 TAMs were pretreated with IL-12 then LPS as compared to LPS alone or IL-12+LPS (Fig. 5E). Overall, IL-12 had no significant impact on the amount TNFα and IL-12 secreted by 4T1 peritoneal TAMs in response to LPS.

Much of IL-12’s impact on myeloid cells has been attributed to its elicitation of IFNγ which further stimulates myeloid cells in an autocrine fashion and activates NK cells and T cells in a paracrine fashion. Macrophages have been shown to produce nitric oxide and upregulate surface molecules associated with antigen presentation in response to IL-12 (169). Both of these effects are
dependent on IFNγ as this phenotype is erased in IFNγ/-/- mice (170, 224). Additionally, IL-12 has been demonstrated to prime macrophages to produce copious amounts of NO in response to LPS (225).

In this study, we demonstrate that IFNγ is important for the IL-12 induced increase in TNFα from 4T1 TIMs and 4T1 peritoneal TAMs in response to LPS (Fig. 13A, Table I, Fig 7A and B). It has been previously shown that IFNγ potentiates the cytoplasmic accumulation of TNFα mRNA (226). It has also been reported that activated monocytes regulate TNFα gene expression both transcriptionally and post-transcriptionally in response to IFNγ (227). A more recent publication showed that IFNγ induced TNFα transcription via transcription factors, IRF-1, IRF-8, and Pu.1 (228). Our data illustrates that 4T1 TIMs from IFNγ-deficient mice produced significantly less TNFα mRNA and protein in response to IL-12 and LPS as compared to wild-type 4T1 TIMs (Fig. 13A) suggesting that the transcriptionally-controlled component of TNFα gene expression in 4T1 TIMs is dependent on IFNγ. Interestingly, although the mRNA levels of TNFα were significantly impacted by the absence of IFNγ, pretreatment of 4T1 TIMs from both wild-type and IFNγ-deficient 4T1 TIMs with IL-12 followed by LPS resulted in a significant increase in TNFα cytokine levels (Fig. 8A, 12A). This presents the possibility that post-transcriptional control of TNFα gene expression in 4T1 TIMs may be IFNγ independent because even in the absence of IFNγ, IL-12 pretreatment of TIMs led to a significant increase in TNFα cytokine production.
Our results show that there was a small but significant difference in the amount of IL-6 gene expression between wild-type and IFNγ-deficient 4T1 TIMs that was not affected by the addition of IL-12 alone (Fig. 13B). We also found that IL-6 gene expression and cytokine production was not significantly different between wild-type and IFNγ-deficient 4T1 TIMs for any of the treatment groups (Fig. 13B, Table I). This suggests that enhancement of IL-6 gene and cytokine expression seen in 4T1 TIMs pretreated with IL-12 followed by LPS is independent of IFNγ. In 4T1 peritoneal TAMs, the absences of IFNγ did not significant impact on the amount IL-6 cytokine produced in response IL-12 and LPS (Fig. 7B). There was, however, a significant increase in IL-6 gene expression in IFNγ deficient 4T1 peritoneal TAMs in response to LPS, IL-12+LPS, and IL-12 pretreatment followed by LPS when compared to WT 4T1 peritoneal TAMs (Fig. 7E). IL-6 is a cytokine with a wide variety of cellular effects and has been implicated in a number of pathological states including cancer. Gene expression of IL-6 is controlled by a number of transcriptional factors, including NF-κβ, C/EβP, AP-1, and Notch signaling proteins (229, 230). Studies have shown that IFNγ enhances the amount of IL-6 secreted in response to LPS in various types of cells (231, 232). Our study shows that IL-12 also increases the amount of IL-6 secreted by 4T1 TIMs in response to LPS in a manner that is independent of IFNγ. This is an important demonstration that IL-12 has a pro-inflammatory impact on TIMs that is not dependent on IFNγ.

We also examined the effect of treatment with IL-12 and LPS on the amount of the anti-inflammatory cytokine, IL-10 secreted from WT 4T1 TIMs and
peritoneal TAMs and IFNγ-deficient 4T1 TIMs and peritoneal TAMs. In our studies, IL-10 cytokine secretion was decreased when 4T1 TIMs from both WT and IFNγ-deficient animals were treated with IL-12 prior to exposure to LPS (Figs. 8E, 12E). We show that there is an overall diminution in the amount of IL-10 secreted by IFNγ-deficient 4T1 TIMs in response to IL-12 and LPS. Our data also demonstrate, however, that the absence of IFNγ did not cause a difference in the amount of IL-10 secreted in any treatment group except for IL-12 alone that was statistically significant when directly comparing WT and IFNγ-deficient 4T1 TIMs (Table I). The amount of IL-10 secreted from TIMs in response to IL-12 was decreased significantly in the absence of IFNγ, although the amount of IL-10 was very low in both groups (Table I). We also observed that pretreatment of 4T1 peritoneal TAMs with IL-12 followed by LPS led to a sharp decrease in the amount of IL-10 as compared to LPS alone and IL-12+LPS (Fig 5H).

It is clear from our data that IFNγ is an important factor in IL-10 gene expression after IL-12 and LPS treatment. When comparing wild-type and IFNγ-deficient 4T1 TIMs directly, there were significant differences in the amount of IL-10 gene expressed in the untreated group and the IL-12 plus LPS group (Fig. 4C). Additionally, there was a difference in IL-10 gene expression in wild-type and IFNγ-deficient 4T1 TIMs in response to LPS alone although it did not reach statistical significance (Fig 4C). We found that IFNγ deficient 4T1 peritoneal TAMs secreted higher amounts of IL-10 and had significantly increased IL-10 gene expression when treated with IL-12 and LPS compared to WT 4T1 peritoneal TAMS (Fig. 7C and F). It has been demonstrated that IL-10 production
is controlled both transcriptionally and post-transcriptionally in manner similar to TNFα by a number of transcription factors, microRNAs, and signaling molecules (233, 234). IFNγ can block IL-10 production through the PI3-kinase-GSK3β-AKT pathway (80). Our results show that even in the absence of IFNγ, IL-12 decreases the amount of IL-10 produced in response to LPS and may affect the post-transcriptional handling of IL-10 in an IFNγ-independent manner. The decreased amount of IL-10 in response to LPS after IL-12 pretreatment may be important in the maintenance or prolongation of IL-12 induced pro-immunological window.

Untreated WT and IFNγ deficient 4T1 TIMs and peritoneal TAMs did not express high levels of IL-10 cytokine that is classically associated with the TIM M2-like phenotype. This observation could be explained by the timing of the measurements of cytokine levels of the untreated group, which was done after overnight incubation. It is reasonable to expect that TIMs lysed immediately after harvest and selection, the IL-10 data may have been different. Several published reports have shown data using TIMs and TAMs from 4T1 and 3LLC tumors with very low levels of IL-10 cytokine expression (180, 235-237). It is hard to determine from in vitro results what would be a functional or physiological relevant amount of IL-10 in vivo. Even so, our data clearly shows that pretreatment of TIMs and TAMs with IL-12 significantly decreases the amount of IL-10 produced in response to the inflammatory stimulus, LPS regardless of the presence of IFNγ. This suggests that IL-12 may prolong the action of other
inflammatory mediators by diminishing the reflexive production of the anti-inflammatory cytokine, IL-10.

We expected that IL-12 would confer similar effects on the cytokine profile of 4T1 TIMs and peritoneal TAMs. We found that although there were some similarities in their response to IL-12 and LPS (i.e. decreased IL-10 after IL-12 pretreatment followed by LPS as compared to LPS alone and IL-12+LPS and IFNγ not required for IL-6 cytokine increases with IL-12 pretreatment), the overall picture was not exactly the same for both TIMs and TAMs. When thinking about possibilities for these differences, two factors come to the forefront. First, all of our experiments were at one set time point (2 hours for mRNA and 18 hours for cytokine data both with 2 hour IL-12 pretreatments). It is possible that this timepoint was optimal for capturing the observation that supported our stated hypothesis for 4T1 TIMs, but was suboptimal for observing similar data in 4T1 peritoneal TAMs. The second factor that may have contributed to differences in the cytokine profile between 4T1 TIMs and 4T1 peritoneal TAMs is the makeup of the myeloid population each group. These pooled samples of CD11b+ cells were not sorted to separate F4/80+ macrophages from Gr1+ monocytic and granulocytic MDSCs or other myeloid cells. We did some simple flow analysis of each group and there were some differences in percentages of different types of cells (data not shown), but a more in depth analysis of the myeloid cell populations of each group of cells would provide more information and might help decipher why there were differences in the response to our stimuli.
We also examined how IL-12 affected the cytokine profile of 4T1 TIMS exposed to tumor derived products. We saw that IL-12 pretreatment significantly enhanced the amount TNFα and IL-6 produced by 4T1 TIMs in response to 4T1 supernatant. IL-12 pretreatment did not affect the high levels of IL-10 secreted from 4T1 TIMs. Assays to determine if myeloid cells pretreated with IL-12 followed by tumor-derived products expressed higher levels of antigen presentation machinery or promoted the activation of T cells would be an interesting follow up experiment. Additionally, determining if IFNγ were necessary for this observation would be an interesting question to pursue. Importantly, this result shows that IL-12 does affect how myeloid cells respond subsequent stimulate and that the tumor microenvironment contains factors with inflammatory potential.

Finally, we examined how several intracellular signaling pathways, including the MAPK pathway, were affected by our treatment of 4T1 TIMs from wild-type and IFNγ-deficient mice with IL-12 and LPS. There is extensive research describing the role of the MAPKs in LPS signaling in macrophages and other myeloid cells (20, 90, 238). Most of the published data concerning IL-12 and the MAPK pathway addresses the pathway’s involvement in IL-12 synthesis in response to inflammatory stimuli such as LPS. Very little work explores how IL-12 stimulation itself affects this pathway. It has been shown in the macrophage cell line, J774E, that IL-12 induces p38 and p44/42 phosphorylation (211). Zhang et al., demonstrated that in T cells, p38 activation is required for IL-12 induced IFNγ production (212). However, this requirement has not been established in
primary macrophages, macrophage cell lines, TIMS, TAMs, or MDSCs. We found that IL-12 alone only induced minimal phosphorylation of the MAPKs wild-type and IFNγ-deficient 4T1 TIMs (Fig. 14). We also observed, as expected, that LPS alone induced phosphorylation of the MAPK pathway in 4T1 TIMs (Fig. 14). Phosphorylation of the MAPK pathway by LPS in 4T1 TIMs was not impacted by the absence of IFNγ (Fig 14). A major observation from these studies is that the increase in MAPK phosphorylation in response to LPS after IL-12 pre-treatment in wild-type 4T1 TIMs was lost in IFNγ-deficient 4T1 TIMs (Fig. 14). Importantly, however, in the IFNγ-deficient 4T1 TIMs, IL-12 pretreatment followed by LPS caused increased phosphorylation of p44/42 at an earlier timepoint when compared to LPS alone (Fig. 14). The early activation of p44/42 in IFNγ-deficient 4T1 TIMs may also contribute to the increases in TNFα and IL-6 with IL-12 pretreatment followed by LPS even in the absence of IFNγ although it would not fully explain the decrease of IL-10 with that same treatment as increased p44/p42 is associated with increased IL-10 expression.
Figure 5. IL-12 modulates the response of 4T1 peritoneal tumor associated myeloid cells to inflammatory stimulus, LPS. CD5+/CD19-/CD11b+ cells isolated from peritoneum of BALB/cJ mice with (4T1 TAMs) or without (control) implanted 4T1 tumors were treated with IL-12, LPS, or IL-12 and LPS for 18 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 18 hours, or left untreated. 100ng/mL of IL-12 and 100ng/mL of LPS were used in all
experiments. TNFα (A, B, C), IL-6 (D, E, F), and IL-10 (G, H, I) protein levels were measured using cytokine bead array. C, F, and I, shows direct comparison between control and 4T1 peritoneal CD5-/CD19-/CD11b+ for each treatment. Figures shown represent the mean ± SEM of triplicate data and are representative of at least three independent experiments with similar results. (*p <0.05, **p<0.001)
Figure 6. 4T1 peritoneal TAMs produce less IFNγ than control peritoneal myeloid cells in the presence of IL-12. For A and B, CD5−/CD19−/CD11b+ cells isolated from peritoneum of BALB/cJ mice with (4T1 TAMs) or without (control) implanted 4T1 tumors were treated with IL-12, LPS, or IL-12+ LPS for 2 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 2 hours, or left untreated. Gene expression was determined by measuring the mRNA levels of IFNγ. In A, control treatment groups are normalized to control untreated peritoneal myeloid cells, while 4T1 treatment groups are normalized to 4T1
untreated TAMs. In B, both control and 4T1 treatments groups are normalized to control untreated peritoneal myeloid cells. For C, CD5−/CD19−/CD11b+ cells isolated from peritoneum of BALB/cJ mice with (4T1 TAMs) or without (control) implanted 4T1 tumors were treated with IL-12, LPS, or IL-12 and LPS for 18 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 18 hours, or left untreated. 100ng/mL of IL-12 and 100ng/mL of LPS were used in all experiments. IFNγ protein levels were measured using cytokine bead array. Figures shown represent the mean ± SEM of triplicate data and are representative of at least three independent experiments with similar results. (*p <0.05, **p<0.001)
Figure 7. The absence of IFNγ affects the response of 4T1 TAMs to IL-12 and LPS. For A, B, and C, CD5−/CD19−/CD11b+ cells isolated from the peritoneum of wild-type and IFNγ-deficient BALB/cJ mice with 4T1 tumors were treated with IL-12, LPS, or IL-12 and LPS for 18 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 18 hours, or left untreated. 100ng/mL of IL-12 and 100ng/mL of LPS were used in all experiments. Cytokine levels were measured using cytokine bead array. For D, E, and F, CD5−/CD19−/CD11b+ cells isolated from the peritoneum of wild-type and IFNγ-deficient BALB/cJ mice with 4T1 tumors were treated with IL-12, LPS, or IL-12+ LPS for 2 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 2 hours, or left untreated. Gene expression was determined by measuring the mRNA levels of each TNFα,
IL-6, and IL-10. All mRNA data shown was normalized to untreated wild-type 4T1 TAMs. Figures shown represent the mean ± SEM of triplicate data and are representative of at least three independent experiments with similar results. (*p <0.05, **p<0.001)
Figure 8. IL-12 modulates the response of 4T1 tumor-infiltrating myeloid cells to inflammatory stimulus, LPS. CD11b+ cells isolated from 4T1 tumors of BALB/cJ mice were treated with IL-12, LPS, or IL-12 and LPS for 18 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 18 hours, or left untreated. 100ng/mL of IL-12 and 100ng/mL of LPS were used in all experiments. TNFα (A), IL-6 (B), and IL-10 (C) protein levels were measured using cytokine bead array. For B, D, E, CD11b+ cells were isolated from 4T1
tumors of BALB/cJ mice and treated with IL-12, LPS, or IL-12+ LPS for 2 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 2 hours, or left untreated. Gene expression was determined by measuring the mRNA levels of TNFα, IL-6, and IL-10 and normalizing to untreated 4T1 CD11b+ cells. Figures shown represent the mean ± SEM of triplicate data and are representative of at least three independent experiments with similar results. (*p <0.05, **p<0.001)
Figure 9. 4T1 TIMs produce IFNγ in response to IL-12 and LPS. On left, CD11b⁺ cells isolated from 4T1 tumors of BALB/cJ mice were treated with IL-12, LPS, or IL-12+ LPS for 2 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 2 hours, or left untreated. Gene expression was determined by measuring the mRNA levels of IFNγ. All treatment groups are normalized to untreated 4T1 TIMs. On CD11b⁺ cells isolated from 4T1 tumors of BALB/cJ mice were treated with IL-12, LPS, or IL-12 and LPS for 18 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 18 hours, or left untreated. 100ng/mL of IL-12 and 100ng/mL of LPS were used in all experiments. IFNγ protein levels were measured using cytokine bead array. Figures shown represent the mean ± SEM of triplicate data and are representative of at least three independent experiments with similar results. (*p <0.05, **p<0.001)
Figure 10

Figure 10. IL-12 increases pro-inflammatory cytokines produced by 4T1 TIMs in response to 4T1 supernatant. CD11b+ cells isolated from 4T1 tumors of BALB/cJ mice were treated with IL-12, LPS, or IL-12 and 4T1 supernatant for 18 hours, pretreated with IL-12 for 2 hours and then treated with 4T1 supernatant for 18 hours, or left untreated. 100ng/mL of IL-12 was used in all experiments. TNFα (A), IL-6 (B), and IL-10 (C) protein levels were measured using cytokine bead array. Figures shown represent the mean ± SEM of triplicate data and are representative of at least two independent experiments with similar results. (*p <0.05, **p<0.001)
Figure 11

Figure 11. Innate immune cells isolated from WT and IFNγ-deficient 4T1 tumor have similar myeloid associated surface markers. Unstimulated CD11b+ cells (A) isolated from 4T1 tumors of WT and Ifng deficient BALB/cJ mice at day 26 were stained with anti-Gr1 (B), anti-F4/80 (C), anti-Ly6C (D), and anti-Ly6G (E). Histograms show fluorescence of each of stained surface marker. Mean fluorescence intensity of surface stain for each group is stated numerically in parenthesis. Curves to the far left of the histogram represent isotype control for WT and IFNγ-deficient samples (as labeled in A). Data shown are representative of 3 independent experiments with similar results.
Figure 12. IL-12 modulates the response of IFNγ-deficient, 4T1 tumor-infiltrating myeloid cells to inflammatory stimulus, LPS. CD11b⁺ cells isolated from 4T1 tumors of Ifng deficient BALB/cJ mice were treated with IL-12, LPS, or IL-12 and LPS for 18 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 18 hours, or left untreated. 100ng/mL of IL-12 and 100ng/mL of LPS were used in all experiments. TNFα (A), IL-6 (B), and IL-10 (C) protein levels were measured using cytokine bead array. For B, D, E, CD11b⁺ cells were isolated from 4T1 tumors of Ifng deficient BALB/cJ mice and treated with IL-12,
LPS, or IL-12+ LPS for 2 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 2 hours, or left untreated. Gene expression was determined by measuring the mRNA levels of TNFα, IL-6, and IL-10 and normalizing to bactin untreated IFNγ-deficient 4T1 CD11b+ cells. Figures shown represent the mean ± SEM of triplicate data and are representative of at least three independent experiments with similar results. (*p <0.05, **p<0.001)
**Figure 13.** IFNγ partially impacts the inflammatory gene profile of 4T1 tumor-infiltrating myeloid cells to IL-12 and LPS. CD11b+ cells were isolated from 4T1 tumors of BALB/cJ mice and Ifng deficient BALB/cJ mice and treated with IL-12, LPS, or IL-12+ LPS for 2 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 2 hours, or left untreated. 100ng/mL of IL-12 and 100ng/mL of LPS were used in all experiments. Gene expression was determined by measuring the mRNA levels of TNFα, IL-6, and IL-10. mRNA fold change averages for each gene were normalized to WT untreated tumor-infiltrating myeloid cells for each treatment. Figures shown represent the mean ± SD of triplicate data and are representative of at least three independent experiments with similar results. (*p <0.05, **p<0.001)
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Table I. IFNγ partially impacts the cytokine response of 4T1 tumor-infiltrating myeloid cells to IL-12 and LPS. CD11b+ cells isolated from 4T1 tumors of BALB/cJ mice and Ifng deficient BALB/cJ mice were treated with IL-12, LPS, or IL-12 and LPS for 18 hours, pretreated with IL-12 for 2 hours and then treated with LPS for 18 hours, or left untreated. 100ng/mL of IL-12 and 100ng/mL of LPS were used in all experiments. TNFα, IL-6, and IL-10 protein levels were measured using cytokine bead array. Average cytokine secretion for each treatment is compared between WT and IFNγ-deficient tumor infiltrating myeloid cells. Figures shown represent the mean ± SD of triplicate data and are representative of at least three independent experiments with similar result.
Figure 14. IL-12 enhances 4T1 tumor-infiltrating myeloid cell activation of MAPK pathway in response to LPS. (A) CD11b+ cells isolated from 4T1 tumors of wild-type BALB/cJ mice were treated with 100ng/mL IL-12 for the indicated time points. After cell lysis, Western blot was performed using antibodies against...
phospho NF-κβ p65 (Ser536), NF-κβ p65, COX-2, NF-κβ p50, and B-actin. CD11b+ cells isolated from 4T1 tumors of wild-type (B) and IFNγ-deficient (C) BALB/cJ mice were treated with IL-12, LPS, or IL-12 and LPS, pretreated with IL-12 for 2 hours and then treated with LPS, or left untreated for indicated time points. 100ng/mL of IL-12 and 100ng/mL of LPS were used in all experiments. After cell lysis, Western blot was performed using antibodies against phospho-p38 (Thr180/Tyr182), total p38, phospho-JNK/SAPK (Thr183/Tyr185), total JNK/SAPK, phospho-p44/42 (Thr202/Tyr204), total p44/42, and B-actin. Data shown are representative of 3 independent experiments with similar results.
CHAPTER IV

CONCLUSIONS AND FUTURE DIRECTION

This work contributes to our knowledge of how IL-12 impacts myeloid cells from the cancer environment. It also demonstrates that IL-12 has IFNγ independent effects on myeloid cells from tumor-bearing animals specifically with regards to IL-6 and IL-10. The relationship between IL-6 and IL-10 is very complicated and interesting. Our data suggests that IL-12, independent of IFNγ, shifts the response to LPS in TIMs and TAMs to more IL-6 and less IL-10. We also show that IL-12 causes increased IL-6 without changes in IL-10 from TIMs in response to tumor derived products. Experiments to determine how IL-12 controls the expression of IL-6 and IL-10 could lead to a better understanding of how these two cytokines contribute to the functional phenotype of myeloid cells.

One of the larger hurdles of cancer immunotherapy is the strong immunosuppression seen in advanced cancer. Early stage cancer are usually easily treated and/or resected, but it is the larger and disseminated tumors that cause the most morbidity and mortality. Agents that combat this profound immunosuppression are attractive immunotherapeutic candidates. IL-12 is a very common immunotherapeutic agent used in clinical trials. Given the broad interest in its anti-cancer effects, it is vital that there is research that focuses on the effect IL-12 on all aspects of the immune system. An area of focus for future work might
include in vivo experiments to determine how myeloid cells within tumors respond to IL-12 in the presence and absence of IFNγ. We know that IFNγ is crucial for IL-12 induced tumor regression (175). This study also showed the IL-12 binds to myeloid populations within the cancer to enhance the effects of adoptively transferred cytotoxic T lymphocytes partly independent of IFNγ. In vivo and ex vivo experiments that observe intrinsic changes in TIMs in response to IL-12 would be helpful in determining the direct impact of IL-12 on TIMs and how that is related to its anti-tumorigenic effects. Our work also shows that IL-12 has IFNγ independent modulatory activity on tumor infiltrating myeloid cells. Our work and the Kerkar paper (175) illustrate the need for more in depth studies on how IL-12 impact TIMs and TAMs to improve cancer immunotherapy. Deciphering IFNγ-independent effects of IL-12 on myeloid cells within cancer may provide new information on how to harness the plasticity of these cells to decrease the immunosuppression that helps maintain cancer.

One of the most interesting aspects of this dissertation is that IL-12 decreased the amount IL-10 secreted in response to LPS in both 4T1 TIMS and peritoneal TAMs. This provides a clue to a possible mechanism by which IL-12 provides a window for an inflammatory stimulus to be more effective. Inflammation is a usually a tightly controlled physiological process where resolution and immunosuppression are important in preventing pathology. In cancer, however, there is pathologic immunosuppression. IL-12 seems to break this immunosuppression and provide a window for other pro-inflammatory stimulus to be more effective. This may be due to direct effects of IL-12 on
myeloid cell IL-10 regulation. Our lab is currently interested in how IL-10 is regulated in macrophages and myeloid cells. Our data show that AMPK is a very important regulator of myeloid cell function and plays a large role in IL-10 production (239, 240). Work that examines the role of IL-12 on IL-10 and AMPK in TIMs may provide critical answers about IL-10 induced immunosuppression and the role of IL-12 in reversing it. Our results are even interesting given work showing that IL-10 has proinflammatory effects in cancer via activation of CD8+ T cells (241, 242). These results reiterate the complexities of immune system alone, as well as, in context cancer. Understanding the multifaceted role of IL-10 in cancer is important in harnessing the power of the immune system to combat cancer.

Research always begets more questions. This work presented here is no exception. Some of these new questions might include: What other signaling pathways are impacted by IL-12 in TIMs? Is the MAPK pathway in TIMs directly involved and necessary for IL-12 induced tumor regression? Are epigenetic changes induced in TIMs after exposure to IL-12 and does this impact the functional phenotype of TIMs? Does IL-12 interaction with the tumor impact the functional phenotype of TIMs? In conclusion, this work provides a small, but important piece of the puzzle as to why and how IL-12 impacts TIMs and TAMs and helps explain its anti-tumorigenic and pro-inflammatory potential.

At the start of this project, although there was great research being done in cancer immunology and immunotherapy, there was also great skepticism about how much cancer immunology and immunotherapy would ultimately
impact cancer patients. The naming of cancer immunotherapy as *Science’s* Breakthrough of the Year for 2013 is proof that investigating the role of the immune system in cancer is a worthwhile pursuit. And while the work done for this project is several large steps removed from clinical application, it is the asking of simple scientific questions like those proposed in this project that lead to cancer immunotherapy being named *Science’s* 2013 Breakthrough of the Year and hopefully to the ultimate goal of eradicating cancer.
REFERENCES


activated protein kinases rather than mitogen-activated protein kinases. 
RING finger protein interacts with the cytoplasmic domain of CD40. 
*J Leukoc Biol.* 67: 2-17.
41. Rothe, M., V. Sarma, V. M. Dixit, and D. V. Goeddel. 1995. TRAF2-
mediated activation of NF-kappa B by TNF receptor 2 and CD40. 
*Science.* 269: 1424-1427.
Induction of the transcription factors NF-kappa B, AP-1 and NF-AT during 
B cell stimulation through the CD40 receptor. *Int Immunol.* 7: 151-161.
Suttles. 2005. TNF receptor-associated factor 6 is an essential mediator of 
CD40-activated proinflammatory pathways in monocytes and 
human leukocyte populations involved in production of interferons alpha 
and gamma. *J Interferon Res.* 1: 233-244.
activates multiple pathways to regulate the expression of the genes for 
major histocompatibility class II I-A beta, tumor necrosis factor and 
complement component C3 in mouse macrophages. *Eur J Immunol.* 19: 
1103-1109.
Identification of interferon-gamma as the lymphokine that activates human 
158: 670-689.
1984. Murine gamma interferon activates the release of a macrophage-
159: 1532-1547.
49. Celada, A., and R. A. Maki. 1991. IFN-gamma induces the expression of 
the genes for MHC class II I-A beta and tumor necrosis factor through a 
and transcriptional activation in response to IFNs and other extracellular 
1992. The structure and function of interferon-gamma receptors. *Int J 
Immunopharmacol.* 14: 413-419.


STAT4 activation by type 1 interferons in the interferon-gamma response to viral infection. *Science.* 297: 2063-2066.


CURRICULUM VITAE

Courtney Mitchell
University of Louisville School of Medicine
Department of Microbiology and Immunology
31. Abraham Flexner Way, Building 55A, Room 313
cjgeor02.louisville.edu, cjgeorge662.gmail.com
(502).852.4622

Education
University of Louisville School of Medicine
MD (May 2014)

University of Louisville School of Medicine
PhD in Microbiology and Immunology (Defense: May 2014, Conferment: August 2014)

Clark Atlanta University,
BS in Biological Sciences (May 2007)
Summa cum laude

Research Experience
Graduate Student, Department of Microbiology and Immunology, University of Louisville School of Medicine, (Fall 2009-present)
Mentors: Robert Stout, PhD and Jill Suttles, PhD
Project title: IL-12 induced immune competence and macrophage functional plasticity and cancer

Student Researcher, Department of Biochemistry and Molecular Biology, University of Louisville School of Medicine, (Summer 2008)
Mentor: Binks Wattenberg, PhD
Project title: “Effects of Sphingosine Kinase Localization on Anchorage Independent Growth”
Investigated the transforming ability of over-expressed sphingosine kinase targeted to various cellular membranes via anchorage independent growth assays.

Student Researcher, Department of Microbiology and Immunology, University of Louisville School of Medicine, (Summer 2007)
Mentor: James Lillard, PhD
Project title: “Transfection and Characterization of CXCR7 expression in Rat Basophilic Leukemia Cells
Rat basophilic leukemia (RBL) cells were transfected with pcDNA-CXCR7, and preliminary FACS was performed with the ultimate goal of identifying biological antagonist for CXCR7.

Student Researcher, Center for Cancer Research and Therapeutic Development, Clark Atlanta University, (Fall 2006-Spring 2007)
Mentor: Shafiq Khan, PhD
Project title: “Biochemical Characterization of Steroidogenesis Inducing Protein (SIP) secreted by ovarian cancer cells”
Worked to characterize steroidogenesis-inducing protein (SIP) from an ovarian cancer cell line to compare with SIP isolated from human follicular fluid

Student Researcher, Pediatric Rheumatology Section, Baylor College of Medicine/Texas Children’s Hospital (Summer 2006)
Mentor: Barry Myones, MD
Project title: “Isolation of the Active Moiety in Lactobacillus casei Cell Wall Extracts Responsible for the Development of Coronary Arteritis”
Analyzed Lactobacillus casei cell wall (LCCW) to isolate and ultimately identify the dynamic element responsible for inducing coronary arteritis using biochemical experimentation

Student Researcher, Center for Biomolecular Science and Engineering, Naval Research Laboratory (Summer 2005)
Mentor: Jinny Lin Liu, PhD
Project entitled “Selection Single Domain Antibodies by Phage Display”
Displayed single domain antibodies from smooth dogfish shark in bacteriophage, transfected E.coli, and then used isolated antibodies to test their affinity for possible biohazardous toxins including cholera, Staphylococcus enterotoxin B, and ricin

Clinical Experience

Resident Physician, Department of Obstetrics and Gynecology, University of Alabama Birmingham (June 2014-June 2018)

Grants
Grant ID: F31CA142317-01
$173,239
07/2009-07/2014
Functional conversion of tumor associated macrophages via cytokine therapy
National Cancer Institute, National Institutes of Health

Publications
IL-12 modification of tumor infiltrating macrophage responsiveness via IFNγ-dependent and independent mechanisms (2013) C. Mitchell, K. Head, J. Suttles, and R.D. Stout. in preparation
Abstracts

Presentations
Poster Presenter, National MD/PhD Conference (2013)
Poster presenter, 3rd Annual National Symposium on Prostate Cancer, CCRTD-CAU (2007)
Poster presenter, Annual Biomedical Research Conference for Minority Students (2005, 2006)

Certifications
Advanced Cardiac Life Support (American Heart Association)
Basic Life Support (American Heart Association)

Honors and Awards
ULSOM Outstanding Student in OB/Gyn Clerkship Award (2014)
John Gibbons Medical Student Award, District V ACOG (2013)
Melvin Denis Travel Award, National MD/PhD Conference (2013)
Doctoral Scholar, Southern Regional Educational Board (2010-present)
Student of Year Award, Department of Biological Sciences, Clark Atlanta University (2007)
Nominated participant, Current Research Topics in Genetics Short Course, National Human Genome Research Institute, National Institutes of Health (2006)
MARC U*STAR Research Program participant, Atlanta University Center (2005-2007)
Provost Scholarship Recipient, Clark Atlanta University (2003-2007)
Academic Dean’s List, Clark Atlanta University (2003-2007)

Memberships/ Committees
University of Louisville School of Medicine Diversity Committee (2012-present)
American College of Obstetricians and Gynecologists, junior fellow (2012-present)
Black Biomedical Graduate Student Organization, president (2010-2011)
Society of Leukocyte Biology, student member (2010-2012)
American Association of Cancer Researchers (2010-2011)
University of Louisville Medical School Student Senate, member (2008-2009)
Student National Medical Association, social chairman (2008-2009)
American College of Physicians, student member (2007-2009)
American Medical Association, student member (2007-2009)
American Medical Women’s Association (2007-2009)

**Community Service**
BBGSO After-School Science Enrichment coordinator (Spring 2010)
Relay for Life Jefferson County, KY (Team captain) (2010)
Black Achievers Louisville volunteer (2007-2008)
Drive Cancer Out volunteer (Spring 2008)