Aryl Hydrocarbon Receptor Antagonism Attenuates Growth Factor Expression, Proliferation, and Migration in Fibroblast-Like Synoviocytes from Patients with Rheumatoid Arthritis

Tejas S. Lahoti, Jarod M. Hughes, Ann Kusnadi, Kaarthik John, Bokai Zhu, Iain A. Murray, Krishne Gowda, Jeffrey M. Peters, Shantu G. Amin, and Gary H. Perdew

Department of Veterinary and Biomedical Sciences, Pennsylvania State University, University Park, Pennsylvania (T.S.L., J.M.H., A.K., B.Z., I.A.M., J.M.P., G.H.P.); DuPont Haskell Global Centers for Health and Environmental Sciences, Newark, Delaware (K.J.); and Department of Pharmacology, Pennsylvania State College of Medicine, Hershey, Pennsylvania (K.G., S.G.A.)

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ABSTRACT

Rheumatoid arthritis (RA) is a chronic autoimmune disease with high morbidity and mortality. Within the inflammatory milieu, resident fibroblast-like synoviocytes (FLS) in the synovial tissue undergo hyperplasia, which leads to joint destruction. Epidemiologic studies and our previous research suggest that activation of the aryl hydrocarbon receptor (AHR) pathway plays an instrumental role in the inflammatory and destructive RA phenotype. In addition, our recent studies implicate the AHR in the regulation of several growth factors, FLS proliferation and migration, along with protease-dependent invasion in FLS from patients with RA (RA-FLS). Treatment with the AHR antagonist GNF351 inhibits cytokine-induced expression of vascular endothelial growth factor-A (VEGF-A), epiregulin, amphiregulin, and basic fibroblast growth factor mRNA through an AHR-dependent mechanism in both RA-FLS and FLS. Secretion of VEGF-A and epiregulin from RA-FLS was also inhibited upon GNF351 treatment. RA-FLS cell migration, along with cytokine-induced RA-FLS cell proliferation, was significantly attenuated by GNF351 exposure. Treatment of RA-FLS with GNF351 mitigated cytokine-mediated expression of matrix metalloproteinase-2 and -9 mRNA and diminished the RA-FLS invasive phenotype. These findings indicate that inhibition of AHR activity may be a viable therapeutic target in amelioration of disease progression in RA by attenuating growth factor release; FLS proliferation, migration, and invasion; and inflammatory activity.

Introduction

Rheumatoid arthritis (RA) is a chronic autoimmune disease with a significant level of morbidity and mortality. Disease progression is characterized primarily by dysregulated proliferation of cells in the synovial lining, such as fibroblast-like synoviocytes (FLS), resulting in hyperplasia, pannus formation, and destruction of associated joint cartilage (Bartok and Firestein, 2010). In the normal synovium, FLS are a highly differentiated unicellular cell type, responsible for providing support, nourishment, and lubrication to the joint tissue. However, in the inflammatory milieu, FLS become hyperplastic, invasive, and highly migratory, reminiscent of tumor cells (Firestein, 1996). FLS hyperplasia serves as a key link between immune cell activity and joint destruction and is considered a hallmark event in RA progression (Qu et al., 1994a). It has been shown that FLS play a constitutive role in the secretion of a number of growth factors, including vascular endothelial growth factors (VEGF), epidermal growth factor (EGF), and fibroblast growth factor (FGF) (Lee et al., 2001; Malemud, 2007; Nah et al., 2010). In turn, growth factor secretion has been shown to activate FLS, leading to hyperplasia and increased angiogenesis, which results in augmented RA severity and progression (Koch, 2000). RA is characterized as a rapidly degenerating disease comprising enhanced FLS invasiveness and migratory behavior. Numerous reports suggest increased expression of

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matrix metalloproteinases (MMPs) and cathepsins in FLS, resulting in destruction of cartilage and bone erosion (Garcia-Vicuna et al., 2004; Giannelli et al., 2004). Each of these events is critical in the progression of RA from local joint inflammation to chronic autoimmune disease. Numerous attempts have been made to target each of these events individually, but few attempts have been made to target multiple events simultaneously.

Apart from genetic factors, clinical and epidemiologic studies suggest that environmental risk factors such as cigarette smoking may contribute to rheumatoid factor seropositivity and bone erosion (Hutchinson and Moots, 2001). Cigarette smoke is a rich source of polycyclic aromatic hydrocarbons, of which many are aryl hydrocarbon receptor (AHR) ligands capable of inducing AHR transcriptional activity through binding to its cognate response element (Marty et al., 2005). The AHR is a ligand-activated transcription factor belonging to the basic helix-loop-helix/Per-ARNT-sim family. Upon ligand-mediated activation, AHR translocates into the nucleus and heterodimerizes with AHR nuclear translocator. This heterodimer then binds to dioxin response elements (DREs), thereby activating the AHR gene battery (Patel et al., 2006). Recently, we have shown that AHR plays an instrumental role in enhancing pleiotropic interleukin-6 (IL6) expression in MCF-7 breast cancer cell lines, leading to enhanced inflammatory signaling (DiNatale et al., 2010b). It has been previously reported that activation of AHR by 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) results in secretion of epiregulin (EREG), an EGF family member and a potent mitogen growth factor capable of enhancing the proliferation of primary mouse keratinocytes (Patel et al., 2006). Amphiregulin (AREG), another member of the EGF family, is secreted by FLS, thereby augmenting the inflammatory response. Previous studies have demonstrated that the AHR can induce amphiregulin expression in the ureteric luminal epithelium (Choi et al., 2006; Yamane et al., 2008). In addition, TCDD has been shown to induce eye vascularization by enhanced production of VEGF through AHR activation (Takeuchi et al., 2009).

We have recently shown that the AHR plays a significant underlying role in regulating proinflammatory IL1B, IL6, and cyclo-oxygenase-2 expression in primary FLS isolated from patients with RA (RA-FLS). In addition, these studies demonstrated that the AHR antagonist GNF351 attenuates cytokine-induced expression of proinflammatory IL1B, IL6, and cyclo-oxygenase-2 levels in an AHR-dependent manner. These studies also established that nuclear translocation of AHR results in binding of the AHR-ARNT heterodimerized to different imperfect DREs present within the IL1B and IL6 genes, thus regulating transcriptional activation (Lahoti et al., 2013). Furthermore, we have determined that constitutive AHR activity in head and neck squamous cell carcinomas (HNSCCs) contributes to their highly invasive and migratory phenotype (DiNatale et al., 2011). Our previous studies also indicate that the AHR antagonist CH223191 inhibits growth factor expression in OSC19 and HNSCC30 cell lines in an AHR-dependent manner (John et al., 2013). Thus, under inflammatory conditions, we hypothesize that constitutive AHR activity plays an important role in growth factor expression, cell proliferation and migration, and invasive phenotype in RA-FLS. Results presented here support this theory in that a potent AHR antagonist, GNF351, attenuates growth factor expression, cytokine-induced proliferation, protease-dependent invasion, and migration in RA-FLS in an AHR-dependent manner. Thus, these data suggest that the attenuation of AHR activity may be a viable therapeutic strategy for the treatment of RA.

### Materials and Methods

N-[2-(3-Indol-3-yl)ethyl]-9-isopropyl-2-(5-methyl-3-pyridyl)purin-6-amine (GNF351) was synthesized as described previously (Smith et al., 2011). TCDD was kindly provided by Dr. Stephen Safe (Texas A&M University, College Station, TX). Human recombinant IL1B was purchased from PeproTech (Rocky Hill, NJ).

#### Cell Culture.

Primary human FLS cells from healthy individuals (N-FLS) and FLS-RA were purchased and maintained in synovocyte growth medium from Cell Application, Inc. (San Diego, CA), unless otherwise indicated.

#### Quantitative Reverse Transcriptase-Polymerase Chain Reaction.

RNA isolation and quantitative reverse transcriptase-polymerase chain reaction (RT-PCR) was performed as previously described (DiNatale et al., 2011).

#### Gene Silencing.

Repression of AHR expression was achieved using small interfering RNA (siRNA) (Dharmacon siRNA, control oligo D-001801-04X, AHR oligo J-004990-07; Pittsburgh, PA). Cells were electroporated with siRNA using the U-020 program of Amaxa nucleofection system (Lonzza, Walkersville, MD) as previously described (Cha et al., 2006; Dinatale and Perdew, 2010). Transfected cells were seeded into six-well plates at 2 × 10⁵ cells/well synovioyte growth media. Cells were cultured for 48 hours post-transfection. Verification of AHR protein ablation was achieved through Western blot analysis.

#### Plasmids.

Minimal HSV-TK-Luc promoter vector was kindly provided by Dr. Jeffrey M. Peters and was digested with KpnI and XhoI. The AREG-HSV-TK-Luc vector construct was made by amplifying 1168 bp of AREG promoter element spanning the region from −384 to −1552 bp with the forward primer 5'-TGC AGG TAC CCA ACA AAT GTG GAA TA-3' and reverse primer 5'-TCA ACT CTA GG CCA AAC AAG GAT AAA GG-3'. The EREG-HSV-TK-Luc vector construct was made by amplifying 1836 bp of EREG promoter element spanning the region from −160 to −1996 bp with the forward primers 5'-TGC AGG TAC CAG CAC TAA CTC GGT GAT-3' and reverse primer 5'-TCA ACT CTG AAA GAA GTG AGC TCA AAT CTG-3'. The EREG-HSV-TK-Luc vector construct was made by amplifying 3053 bp of EREG-A promoter element spanning the region from +1043 to −2010 bp with the forward primers 5'-TGC AGG TAC CAG CAG CAC ACA AGG AG-3' and reverse primer 5'-TCA ACT CTG AGC ACC CAA GAC AGC AG-3'. Each primer contained the appropriate restriction enzyme site for insertion into the vector.

#### Transient Transfection and Luciferase Assay.

COS-1 cells were maintained in α-minimum essential media (MEM) with 10% fetal bovine serum (FBS), 50 units/ml penicillin, and 50 µg/ml streptomycin. Upon reaching ~80% confluence, COS-1 cells were transiently transfected with pSil/β-Gal (100 ng/well), pcDNA3-hAHR (500 ng/well), and either VEGF-A-HSV-TK-Luc or EREG-HSV-TK-Luc or AREG-HSV-TK-Luc plasmids (100 ng/well) using Lipofectamine 2000 (500 ng/well), and either VEGF-A-HSV-TK-Luc or EREG-HSV-TK-Luc or AREG-HSV-TK-Luc plasmids (100 ng/well) using Lipofectamine Plus (Invitrogen, Carlsbad, CA), according to the manufacturer's protocols. After 24 hours, cells were treated with vehicle [dimethylsulfoxide (DMSO)] or TCDD (10 nM) for 4 hours. Luciferase activity was determined using Luciferase Assay System Substrate (Promega, Madison, WI). Transfection efficiency and data normalization were performed using β-galactosidase activity.

#### Enzyme-Linked Immunosorbert Assay.

For the measurement of VEGF-A and EREG levels, RA-FLS cells were seeded at 2 million cells per ml and pretreated with GNF351 for 1 hour, followed by stimulation with IL1B. Cell supernatants were centrifuged to avoid any cell debris prior to enzyme-linked immunosorbent assay (ELISA) analysis. To determine the levels of growth factor secretion, VEGF-A (Abnova, Taipei, Taiwan) and EREG (Usen Life Science, Wuhan, China) ELISAs were performed per manufacturers' instructions.
Bromodeoxyuridine Staining. RA-FLS cells were plated in chamber slides and treated with GNF351 in the presence or absence of IL1B for 48 hours. Cells were then treated with 10 μg/ml bromodeoxyuridine (BrdU) for 2 hours. Cells were subsequently fixed with 4% paraformaldehyde for 15 minutes and permeabilized with 0.3% Triton X-100 at room temperature for 15 minutes, then treated with 2 N HCl at 37°C for 30 minutes, followed by neutralization with 0.2 M borate buffer (pH 8.3). Cells were blocked with 2% normal mouse serum in phosphate-buffered saline (PBS) with 0.2% Tween 20 (PBST) for 30 minutes and then incubated with Anti-BrdU fluorescein (Abcam, Cambridge, MA) overnight at 4°C in PBS with 0.1% Triton X-100 and 0.5% bovine serum albumin (BSA). The next day cells were washed three times with PBST, followed by nuclei staining with Hoechst solution (final concentration, 1 μg/ml) for 20 minutes. Fixed cells were again washed with PBST, and the slide was mounted with ProLong Gold Antifade reagent (Life Technologies, Grand Island, NY). Specimens were examined on a fluorescence microscope. Data were plotted by counting the BrdU-positive cells from eight different fields and normalized to total number of cells in each field per individual treatment.

Ki-67 Staining. RA-FLS cells were treated as described earlier for BrdU staining in chamber slides and fixed with 2% paraformaldehyde for 15 minutes, followed by permeabilization with 100% methanol for 10 minutes at −20°C. Upon methanol permeabilization, cells were blocked with 2% normal mouse serum in PBST for 30 minutes. Ki-67 antibody conjugated with Alexa Fluor 555 (BD Pharmingen, San Jose, CA) was placed in PBS with 0.1% Triton X-100 and 0.5% BSA. Cells were incubated overnight at 4°C and washed three times with PBST. Nuclei were stained with Hoechst solution. The cells were examined on a fluorescence microscope.

Zymography. Cell culture supernatants (50-μg protein) were mixed with 2× gel loading buffer (3 M Tris HCl, pH 8.45, 25% glycerol, 0.8% SDS, 0.1% Coomassie Blue, and 0.1% phenol red) and then resolved under nonreducing conditions by 8% SDS-PAGE embedded with 3 mg/ml gelatin (Sigma-Aldrich, St. Louis, MO). Gels were rinsed three times in 2.5% Triton X-100 for a total of 30 minutes at room temperature, followed by three times with water for 15 minutes at room temperature. The gels were incubated in substrate buffer (50 mM Tris HCl, pH 7.45, 10 mM CaCl2) overnight at 4°C and washed three times with PBST. Gels were stained with Coomassie Blue. Areas of gelatinolytic activity were visualized as transparent bands.

**Results**

GNF351 Inhibits IL1B-Induced Growth Factor Expression in FLS. An inflamed synovium has been shown to contain elevated levels of growth factors, which in part induce FLS hyperplasia. It is proposed that secretion of inflammatory cytokines into synovial fluid results in FLS proliferation, which in turn activates FLS along with other cell types within the synovial lining, leading to further release of inflammatory cytokines and growth factors via autocrine and paracrine loops. Our recent studies in HNSCC cell lines have revealed that the AHR plays a central role in the constitutive regulation of a number of inflammatory and growth factor genes (John et al., 2013). Thus, we wished to determine the effect of constitutive AHR activity and AHR antagonist treatment on FLS growth factor expression in an inflammatory environment. N-FLS and RA-FLS cells were treated with GNF351 for 1 hour, followed by cytokine challenge with IL1B for 4 hours. The results indicate that pretreatment with GNF351 significantly inhibits IL1B-mediated enhanced expression of EREG, AREG, VEGF-A, and FGF-2 mRNA in N-FLS (Fig. 1A) and RA-FLS (Fig. 1B). GNF351 exhibited similar ability to inhibit growth factor expression in RA-FLS from several patients with RA (data not shown). This is consistent with the previous results obtained from five patients upon examining the ability of GNF351 to attenuate IL1B expression (Lahoti et al., 2013). It is important to note that in this previous report we demonstrated that long-term exposure to 1 μM GNF351 failed to exhibit any observable cellular toxicity (Lahoti et al., 2013).

**GNF351-Mediated Repression of Growth Factor Expression Is AHR-Dependent in FLS.** We previously reported that GNF351-mediated repression of proinflammatory gene expression in RA-FLS cells is dependent on AHR expression (Lahoti et al., 2013). A similar approach was used to demonstrate that attenuation of growth factor expression following GNF351 treatment in N-FLS and RA-FLS is also AHR-dependent. Repression of AHR expression was achieved through transient siRNA-mediated protein ablation, which led to an ~75% decrease in AHR protein expression in N-FLS and RA-FLS. This resulted in both cell types being unresponsive to GNF351 treatment upon IL1B-mediated induction of EREG, AREG, VEGF-A, and FGF-2 expression (Fig. 2, A and B). Furthermore, ablation of AHR protein levels in both N-FLS and RA-FLS resulted in FLS becoming more refractory to the ability of IL1B to induce growth factor expression. These results support the concept that the AHR is involved in contributing to the level of IL1B-mediated induction of several growth factors in FLS and demonstrate that the effect of GNF351 is dependent on AHR expression.

**The Promoters of Growth Factor Genes Contain Functional DREs.** Putative DRE-like sequences were identified in the promoter regions of AREG, EREG, and VEGF (Supplemental Fig. 1). This observation led to the hypothesis that these genes are directly regulated by the AHR through the presence of functional DRE-like elements. Promoter fragments containing DRE-like elements were cloned into a minimal HSV-TK-Luc promoter vector. Increased transcriptional activity was observed in TCDD-treated COS-1 cells.
which were transfected with pcDNA3-hAHR and cotransfected with either VEGF-A-HSV-TK-Luc, EREG-HSV-TK-Luc, or AREG-HSV-TK-Luc (Fig. 3A). TCDD-dependent activation of AHR in COS-1 cells led to enhanced transcriptional activity for each growth factor tested. 

**Antagonism of the AHR Attenuates Secretion of VEGF-A and EREG.** Growth factor release has been associated with enhanced proliferation of synoviocytes, along with angiogenesis in synovial tissue. Therefore, synoviocytes from RA patients were cultured for 48 hours in the presence of IL1B, and the ability of GNF351 cotreatment to attenuate VEGF or EREG secretion was examined. The results demonstrate that GNF351 can inhibit IL1B-induced secretion of VEGF-A and EREG from RA-FLS cells (Fig. 3B).

**GNF351 Inhibits IL1B-Induced RA-FLS Proliferation.** Synovial lining hyperplasia, a hallmark event in RA, is primarily associated with uncontrolled FLS proliferation (Qu et al., 1994a). Proinflammatory cytokines can induce FLS transformation in RA, which in turn causes a high degree of FLS proliferation in vivo and in cell culture models (Lacey et al., 2003; Leech et al., 2003). AHR activation has been shown to directly regulate cell cycle progression and tumor cell proliferation (Shimba et al., 2002). Thus, to study the effect of the AHR antagonist GNF351 on cytokine-mediated RA-FLS proliferation, RA-FLS cells were subjected to BrdU and Ki-67 immunofluorescent staining (Fig. 4). The data indicate that IL1B significantly stimulates RA-FLS DNA synthesis. In contrast, treatment of RA-FLS with GNF351 significantly repressed cytokine-induced RA-FLS DNA synthesis, suggesting a key role for the AHR in cytokine-induced FLS hyperplasia.

**GNF351 Inhibits FLS Migration.** Elevated levels of chemokines and cytokines in RA synovium enhance FLS migration, contributing to RA progression both within one joint and through migration to other joints. To illustrate the effect of the AHR antagonist GNF351 on FLS migration, real-time analysis of cell migration with the xCELLigence System was performed. RA-FLS cells treated with vehicle or GNF351 did not migrate in the absence of serum. The presence of serum triggered a chemotactic response demonstrated by the marked (~50-fold) increase in RA-FLS migration following treatment with vehicle; in contrast, GNF351 treatment significantly attenuated RA-FLS migration (Fig. 5A). The effect of GNF351 on RA-FLS migration was also examined in a scratch assay. Cell migration was evident in the scratch assay, with significant migration observed after 10 hours post-scratch (data not shown) and marked RA-FLS migration across the scratch observed after 16 hours (Fig. 5B). However, the presence of GNF351 significantly attenuated cell migration across the scratch. It is important to note that the effects were independent of cell proliferation, as primary RA-FLS require ~36 hours for one replication cycle. These results suggest that the presence of an AHR antagonist attenuates chemoattractant-induced FLS migratory behavior.

**GNF351 Inhibits Cytokine-Induced MMP-2 and -9 Expression in RA-FLS.** Inflamed synovial tissue has been shown to be highly invasive, leading to the progressive destruction of cartilage and bone. In addition, RA-FLS have the ability to secrete zinc-dependent proteases such as MMPs, which promote degradation of the extracellular matrix proteins in joint tissue and bone (Murphy and Nagase, 2008). Recent reports
GNF351-mediated repression of growth factor expression is AHR-dependent in primary FLS. Repression of AHR expression was performed using AHR-specific siRNA in primary FLS from healthy donors (A) or from patients with RA (B). A total of $2 \times 10^6$ cells were used per gene knockdown sample. The repression of AHR protein expression was confirmed using Western blot analysis after 48 hours of siRNA transfection. Upon confirmation, primary N-FLS transfected with siRNA for 48 hours were pretreated with 500 nM GNF351 for 1 hour, followed by 10 ng/ml IL1B challenge for 4 hours. The levels of EREG, AREG, VEGF-A, and FGF-2 mRNA were determined by quantitative RT-PCR analysis. Data points and bars represent mean $\pm$ S.E. of three independent determinations. Data were analyzed using one-way ANOVA followed by Tukey's multiple-comparisons test. $^* P < 0.05; ^{**} P < 0.01; ^{***} P < 0.001$. Alphabetical characters indicate statistical comparisons between two groups.
suggest that AHR plays an active role in regulating MMP expression during normal development, as well as in numerous cancers (Hillegass et al., 2006; Ishida et al., 2010). Thus, to determine the effects of the AHR antagonist GNF351 on the expression of MMPs, RA-FLS were treated with GNF351 in the presence or absence of proinflammatory IL1B. The results indicate that IL1B significantly induced mRNA expression of MMP-2 and -9, while GNF351 pre-treatment attenuated IL1B-mediated MMP-2 and -9 induction (Fig. 6A; Supplemental Fig. 1). The functional significance of this repression by GNF351 was validated by gelatin zymography, in that IL1B-induced pro-MMP-2- and -9-mediated gelatin degradation was significantly mitigated by GNF351 treatment (Fig. 6B; original color image: Supplemental Fig. 2).

**Discussion**

Although the etiology of RA has not been firmly established, it is now known that T-cell-mediated inflammatory responses lead to an inflamed synovium, resulting in release of cytokines and chemokines and subsequent activation and recruitment of peripheral mononuclear cells (Cope, 2008). These events lead to activation of resident FLS and fibroblast-like macrophages. Increasing levels of cytokines and chemokines in the synovium lead to activation of FLS, which undergo hyperplasia, a hallmark event in RA. Activated FLS also migrate throughout the joint and even invade other joints, leading to extensive cartilage and bone deterioration (Firestein, 1996). FLS have also been shown to secrete multiple cytokines and growth factors that in part activate an autocrine loop, resulting in further FLS hyperplasia (Qu et al., 1994b; Afuwape et al., 2002). Although the etiology of RA is complex, both genetic and environmental factors have been identified. There is an increasing body of epidemiologic research pointing to smoking as a risk factor for RA. The fact that tobacco smoke is a rich source of planar polycyclic aromatic hydrocarbons, many of which are AHR agonists (e.g., benzo[a]pyrene), suggests that activation of the AHR may play a significant role in disease progression (Baka et al., 2009). Furthermore, smoking has been shown to modulate the immune system by altering Th17 cell–mediated responses (Onozaki, 2009;
Marshall and Kerkvliet, 2010). Thus, we hypothesized that AHR antagonism may serve as a viable therapeutic target for RA treatment.

GNF351 is a complete antagonist that inhibits both AHR-mediated, DRE-dependent activity induced by agonists as well as selective AHR modulator-mediated, DRE-independent AHR signaling (Nah et al., 2010; Zhao et al., 2010; Smith et al., 2011). GNF351 has been shown to inhibit nuclear translocation of the AHR in an immortalized FLS line (K4IM), thus inhibiting DRE-mediated transcription of inflammatory cytokines such as IL1B and IL6 (Lahoti et al., 2013). Other possible therapeutic activities attributed to AHR antagonists include SR1, a potent AHR antagonist used in hematopoietic stem cell expansion ex vivo, and the AHR antagonist 6,2',4',9-trimethoxyflavone, which inhibits expression of IL6 in multiple HNSCC cell lines in the presence of inflammatory signaling. These examples suggest that AHR antagonists exhibit therapeutic potential for the treatment of a variety of diseases (Boitano et al., 2010; DiNatale et al., 2011).

Proinflammatory cytokines such as IL1B, tumor necrosis factor-α, and IL6 regulate growth factor expression and subsequent secretion from RA-FLS. The data presented here suggest that GNF351 pretreatment of FLS from healthy or RA-positive individuals costimulated with proinflammatory IL1B can also attenuate cytokine-induced expression of multiple growth factors, such as VEGF-A, EREG, AREG, and FGF-2. We have recently demonstrated that there are multiple imperfect DREs present in the promoter regions of FGF-2, EREG, and AREG (John et al., 2013). Thus, we believe that the effects elicited by GNF351 likely occur through inhibition of AHR recruitment to DREs present in the promoter regions of these growth factors and are independent of extracellular signal-regulated kinase and nuclear factor-κB activation pathways (Lahoti et al., 2013). Consequently, our data now provide new insight in that AHR antagonism not only mitigates cytokine release but also inhibits growth factor expression.

Synovial angiogenesis plays an instrumental role in the invasive and destructive synovial tissue phenotype. The release of VEGF and other growth factors enhances angiogenesis in inflamed synovial tissue, and thus elevated serum growth factor levels can serve as an indicator of RA severity. The fact that the AHR actively enhances IL1B, IL6, and VEGF secretion in RA-FLS, coupled with the fact that a recognized treatment for RA is an anti-IL6 receptor monoclonal antibody that lowers serum VEGF levels, suggests that using an AHR antagonist as a therapeutic treatment is worthy of further consideration (Lee et al., 2001; Nakahara et al., 2003; Roman et al., 2009; DiNatale et al., 2010b). High levels of cytokines and chemokines in synovial fluid transform FLS to undergo proliferation, forming a pannus tissue, which further secretes more cytokines and growth factors (Firestein, 1996). Activation of the AHR has been shown to induce cell proliferation, linking inhibitory effects of AHR antagonists in cytokine-induced RA-FLS.
proliferation (Shimba et al., 2002; Chang et al., 2007). Transformed FLS also undergo migration and can induce protease-dependent invasion of surrounding bone and tissue cartilage. We have previously shown in HNSCC that treatment with the AHR antagonists 2',4'-trimethoxyflavone and GNF351 or ablation of AHR protein levels results in cells that are highly refractory to cellular migration (DiNatale et al., 2012). This further supports our current findings, indicating a regulatory role for the AHR in RA-FLS migration. FLS derived from inflamed RA synovium possess an invasive phenotype, in contrast to FLS isolated from healthy individuals or osteoarthritis patients (Zvaifler and Firestein, 1994). FLS-RA have also shown the ability to invade normal human cartilage engrafted in SCID mice (Muller-Ladner et al., 1996). Activation of the AHR pathway by an exogenous ligand like TCDD in lung airway epithelial cells, normal human keratinocytes and melanocytes, and the melanoma cell line A2058 induces expression of MMP-2 and -9 (gelatinase A and B) mRNA expression; however, this induction was significantly attenuated by GNF351 treatment. Although MMP-2 and -9 are gelatinase-specific, numerous reports suggest that increasing levels of MMP-2 and -9 in synovial fluid can be used as a parameter of RA progression and these MMPs can degrade fibrillar collagen molecules (Ahrens et al., 1996; Yoshihara et al., 2000; Itoh et al., 2002).

The ability of AHR to participate in growth factor and cytokine expression in synoviocytes from normal individuals firmly supports the notion that AHR activation may play a significant role in the development and progression of RA. Furthermore, the fact that RA-FLS and N-FLS were cultured in medium that is largely devoid of exogenous AHR ligands makes the data obtained here even more remarkable. Taken together, this strongly suggests that AHR activation occurs in synoviocytes either through production of endogenous ligands or through unliganded AHR activation. It is likely that there are a number of potent AHR endogenous ligands present in

Fig. 5. GNF351 inhibits migration of primary FLS from patients who tested positive for RA. (A) Primary RA-FLS were treated for 12 hours with GNF351 or vehicle alone; after 8 hours, cells were serum-starved for 4 hours. Cells were then plated at 5000 cells per top well of a CIM-Plate 16 in serum-free media with or without GNF351. The bottom wells were filled with either DMEM or DMEM with serum as a chemoattractant. (B) Primary RA-FLS were plated in six-well plates. Cells were treated for 12 hours with 500 nM GNF351 or vehicle alone in triplicate. After 12 hours, a scratch was made in the middle of the well with a sterile p200 pipette tip. At 1 hour post-scratch, cells were retreated with GNF351 or vehicle alone every 12 hours for a total period of 16 hours; wells were photographed immediately after scratch and 16 hours post-scratch. Data points and bars represent mean ± S.E. of three independent determinations. Data were analyzed using one-way ANOVA followed by Tukey's multiple-comparisons test. ***P < 0.001. Alphabetical characters indicate statistical comparisons between two groups.
humans, including indoxyl sulfate and kynurenic acid, especially during disease progression (DiNatale et al., 2010a; Schroeder et al., 2010). In addition, we have previously demonstrated that kynurenic acid increases IL6 production in the presence of IL1B, which is dramatically inhibited by GNF351 treatment. It is noteworthy that kynurenic acid has been detected in synovial fluid from patients with RA at concentrations that are within the range capable of inducing significant AHR activation (Lahoti et al., 2013). In addition, exogenous AHR ligands can be derived from the diet, such as indole-3-carbinol, a breakdown product of glucobrassicin found in cruciferous vegetables, further adding to the total pool of circulating AHR ligands. Our preliminary data suggest that synovial fluid from patients with RA has significant potential to activate the AHR (unpublished data). The enhanced AHR expression observed in clinical samples further supports an important role for the AHR in RA (Lahoti et al., 2013). Enhanced AHR expression has also been seen in tumor cells when compared with non-neoplastic parental control cells (DiNatale et al., 2012). Thus, it is anticipated that the AHR would play an even greater role in synoviocyte growth factor and cytokine expression in vivo because of persistent ligand activation, compared with the in vitro data presented here and in our previous report (Lahoti et al., 2013). Overall, our findings indicate that the AHR provides a viable therapeutic target for amelioration of disease progression in RA by inhibiting growth factor release, RA-FLS proliferation and migration, and as previously reported, cytokine expression (DiNatale et al., 2012). Future studies are needed to establish whether AHR antagonism might be used as a treatment strategy.

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**Authorship Contributions**

Participated in research design: Perdew, Lahoti.

Conducted experiments: Lahoti, Hughes, Kusnadi, John, Zhu, Murray.

Conducted new reagents or analytic tools: Gowda, Amin, Peters.

Performed data analysis: Lahoti, Perdew, Peters.

Wrote or contributed to the writing of the manuscript: Lahoti, Perdew.

**References**


Address correspondence to: Dr. Gary H. Perdew, Center for Molecular Toxicology & Carcinogenesis, Department of Veterinary and Biomedical Sciences, Pennsylvania State University, 309A Life Sciences Building, University Park, PA 16802. E-mail: ghp2@psu.edu