Disruption of Angiogenesis by Anthocyanin-Rich Extracts of *Hibiscus sabdariffa*

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Abstract—Abnormal vessel formations contribute to the progression of specific angiogenic diseases including age-related macular degeneration. Adequate vessel growth and maintenance represent the coordinated process of endothelial cell proliferation, matrix remodeling, and differentiation. However, the molecular mechanism of the proper balance between angiogenic activators and inhibitors remains elusive. In addition, quantitative analysis of vessel formation has been challenging due to complex angiogenic morphology. We hypothesized that conjugated double bond containing-natural products, including anthocyanin extracts from *Hibiscus sabdariffa*, may control the proper angiogenesis. The current study was designed to determine whether natural molecules from African plant library modulate angiogenesis. Further, we questioned how the proper balance of anti- or pro-angiogenic signaling can be obtained in the vascular microenvironment by treating anthocyanin or fatty acids using chick choioallantoic membrane angiogenesis model in ovo. The angiogenic morphology was analyzed systematically by measuring twenty one angiogenic indexes using Angiogenic Analyzer software. Quantitative analysis showed that anthocyanin is an angiogenic modulator that can be used to treat uncontrolled neovascular-related diseases, including age-related macular degeneration.  

Index Terms—age-related macular degeneration, angiogenesis, anthocyanin, chick embryo, choioalantoic membrane, fatty acid, *Hibiscus sabdariffa*, molecular modeling, VEGFR, vessel formation

1 INTRODUCTION

BALANCED blood vessel formation corresponds to the assembly of angioblasts, vessel sprouting, recruiting pericytes and endothelial smooth muscle cells to form tubes, branches, and capillary blood vessels from the existing vascular network [1–4]. Uncontrolled angiogenic reactions are the major pathological components of the late stage of wet age-related macular degeneration (AMD) when abnormal blood vessels develop toward into the macular. Pathological angiogenesis is also seen in other inflammatory diseases, ischemic conditions, diabetic neuropathy, and cancer, thus disruption of angiogenesis has become a promising target for therapies to stop diseases that thrive on angiogenesis. However, the molecular mechanism of the proper balance between angiogenic activators and inhibitors remains unknown. In addition, systematic analysis of angiogenesis has been challenging due to complex vessel morphology that includes vessel length, branches, nodes, and segments.

Here, we propose that natural compounds containing aromatic rings with a carbohydrate moiety in plant library, may regulate blood vessel formation through the inhibition of angiogenesis. The current study was designed to determine the time and dose-dependent modulation of angiogenesis by anthocyanin and how the proper analysis could be performed using systematic calculations.

Previously, our proteomic data demonstrated that early signaling molecules, including erythropoietin, prohibitin, PP2A, vimentin, HIF1, JAK2, and nitric oxide, are involved in apoptosis and angiogenesis in the retina and the RPE, and imbalance of oxidative environment may lead to the initiation of AMD [5–11]. Here, we aimed to use specific natural compound library to regulate the early signaling targets to modulate blood vessel formation processes. The therapeutic role of *Hibiscus sabdariffa* calyx was examined and its extracts linked to its phytochemical constituents including anthocyanin (>100 mg cyanidine rutinoside/100 g p lants) were determined [12,13]. We established 21 angiogenic index, including vessel length, branches, nodes and segments, using Angiogenesis Analyzer software. Quantitative analysis showed that anthocyanin inhibited 50-80 % of blood vessels formation. As angiogenic activators, selected fatty acid mixtures that include oleic (18:1), linoleic (18:2, w6), linolenic (18:3, w3), palmitic (16:0), and palmitoleic (16:1) acid, significantly stimulated angiogen-
esis by 20-50%. Our data suggest that natural chromophore from edible plants library may regulate the vessel formation in time- and dose-dependent manner. Protein-ligand docking analysis using SwissDock software proposed that hibiscetin from Hibiscus may bind to the active site of Vascular endothelial growth factor receptor 2 (VEGFR2) with the Asp1046, Cys1045, Cys919, and Glu917 showing ΔG=-8.62 kcal/mol of binding energy. The current study implies that anthocyanin rich extract of H. sabdariffa has an anti-angiogenic effect in ovo and may be a potential candidate for the treatment of angiogenic diseases.

2 MATERIALS AND METHODS

2.1 Methods
We followed the NIH Guide and the Association for Research In Vision and Ophthalmology (ARVO) statement for in vivo experiments. Avian embryos are not considered live animals under PHS policy. The use of chicken embryos at gestation day 12 and younger does not require an ACUC Animal Use Protocol Application. Chick embryos younger than embryonic day 13 (E13) are assumed unable to experience pain. Therefore, E13 and younger embryos were euthanized by hyperthermia, typically conducted by placing the eggs in a -20°C freezer or < 4°C for 4 hours. Embryonic death was confirmed by decapitation, membrane disruption, or maceration. Chick embryos between E13 – E17 can experience pain and were euthanized by cervical dislocation. All experiments were repeated with many biological samples (n >3) with technical triplicate (n > 9). Stat View software was used for statistical analysis. Statistical significance was analyzed using unpaired Students’ test or variance (ANOVA) when appropriate. The significance level P < 0.05 is considered as statistically significant.

2.2 Collection of Plant Materials
Air dried anthocyanin-containing plants library, including Hibiscus sabdariffa calyx, Moringa oleifera, Hibiscus asper, Vernononia amygdalina, Elaeis guineensis, Manihot esculenta, Chrysanthemum morifolium, Sesamum radiatum, Balanite aegyptica, Talinum triangulare and Telfairia occidentalis was obtained locally from Yola, Nigeria. It was taxonomically identified and authenticated using three different African plant databases (http://www.gbif.org/resource/81181; http://posa.sanbi.org/intro_precis.php; http://www.africanplants.senckenberg.de/root/index.php).

2.3 Extraction of Anthocyanin
H. sabdariffa calyx extracts were obtained by soaking 100 g of the powder in 1.8 L of distilled water for 12 hours, followed by filtering the mixture using filter paper (pore size 11 μm). The filtrate was collected and the residue rinsed with 200 mL of distilled water. The filtrate was lyophilized and used for the current study. Anthocyanin concentrations, both total and monomeric, were determined quantitatively by pH differential method. Anthocyanin content was calculated as mg cyanidin-glucoside equivalent/100 g of solid sample (ε = 28840 M⁻¹•cm⁻¹) [12,13]. Fatty acids (unsaturated fatty acid: saturated fatty acids = 85:15, oleic/linoleic/linolenic/palmotilic/palmitic/stearic acid) were tested as pro-angiogenic molecules [14,15].

2.4 Inoculation of Chick Embryo with Anthocyanin
Freshly prepared H. sabdariffa calyx extracts (dissolved in Ringer’s solution or PBS or ddH₂O) 14 µg or 58 µg concentrations were injected in fertilized eggs (average 38 g, chick embryo, Gallus domesticus) at 0, 2, 4, 7 , and 9 days (pre-incubation) and were incubated at 37.5 °C for additional 4, 5, 7, 15 days (post-incubation) in humidified incubator. Angiogenic indexes such as vessel length, branches, nodes and segments were determined upon treatment with anthocyanin. The experiments were repeated 20 times to determine time and dose-dependent vessel formation on anthocyanin treatment. Each experiment includes 7 - 30 chick embryos/set/week and a total of 212 chick embryos were used and 345 vessel images were obtained. Pre-incubation day ED = 0-4 days and Post-incubation day ED = 4-10 days were mainly tested against control sample that contains the same amount of Ringer solution.

2.5 Angiogenic Analysis of Chick Embryo Chorioalatoic Membrane
Twenty one angiogenic parameters were tested using Angiogenesis Analyzer macro connected Image J software (Gilles Carpentier, http://image.bio.methods.free.fr/ImageJ?Angiogenesis-Analyzer-for-ImageJ). Vessel formation of anthocyanin-treated chick embryo with pre-incubation (0-2 days) and post-incubation (4-7 days, anthocyanin + = 14 µg, and anthocyanin ++ = 58 µg/chick embryo) was analyzed. Angiogenic Analysis showing extremities (A1-B1); nodes identified as 3 neighbor (A2-B2); twig (C1, D1), segment (C2, D2) delimited by two junctions (C3, D3) and branch (C4, D4). E shows a junction implicated only in branch (E1) and master junctions like E2 delimiting master segments (E3). F represents the master tree composed from master segments associated by master junctions delimiting the meshes (F1). Two close master junctions can be fused into a unique master junction (F2) and the underlying segment (F3).

2.6 Molecular Modeling
A 3D model of the protein was built using the 3D structure VEGFR2 as a template and selected ligands, including malvidin 3-galactoside, cyanidin, cyaniding 6-glucoside, and hibiscetin [16-18]. A random selected non-VEGFR2 binding mole-
cules, including alanine, D-glucose, and oxalic acid were used as negative controls (data not shown).

Vascular endothelial growth factor receptor 2 (VEGFR2) was obtained from the Protein Data Bank. Protein-ligand docking server, including SwissDock and GOLD (version 5.0), was used for ligand-protein interaction studies (http://www.swissdock.ch/docking?). The active site of VEGFR2 and protein structure were examined among seven X-ray diffraction crystal structures (PDB ID = 1Y6A, 1Y6B, 1YWN, 2OH4, 2P2H, 2P21, 2QUS). The conserved amino acid residues presenting the VEGFR2 a citive site were L840, V848, C866, K868, E885, V899, V916, E917, F918, C919, K920, G922, L1035, C1045, D1046 and F1047. SwissDock predicted the molecular interaction between VEGFR2 and selected anthocyanin molecules, including hibiscetin. SwissDock contains a database of proteins and ligand structures. SwissDock generated many binding modes using EADock software as local docking in the box as well as blind docking in the vicinity of all VEGFR2 cavities. At the same time, binding energies were calculated and the most favorable energies were evaluated to show the most favorable clusters. 257 bindings were calculated with clusters (0-37), elements (0-15), FullFitness (-1588.73 to -1567.50 kcal/mol), binding energy ΔG (-8.42 to -6.20 kcal/mol). SwissDock used a hybrid evolutionary algorithm with two fitness functions, with the CHARMM package for energy calculations.

3 RESULT AND DISCUSSION

3.1 Anti-angiogenic Effect of Anthocyanin-Rich Extract of Hibiscus sabdariffa

First, we tested the following air dried anthocyanin-containing plants as an initial library, including Hibiscus sabdariffa calyx, Moringa oleifera, Habiscus asper, Vernonia amygdalina, Elaeis guineensis, Manihot esculenta, Chrysanthemum morifolium, Sesamum radiatum, Balanite aegyptica, Talinum triangulare and Telfairia occidentalis, obtained locally in Yola, Nigeria (Figure 1). Freshly prepared Hibiscus extracts (dissolved in Ringer’s solution or ddH2O) were injected in fertilized eggs (Gallus domesticus) at 0, 2, 4, 7, and 9 days and were incubated at 37.5 °C for additional 4, 5, 7, 15 days [19]. The altered angiogenesis upon anthocyanin treatment was analyzed by angioenic indexes. Pre-incubation embryonic day (ED) = 0, 2, 4, 9 d ays and post-incubation embryonic day ED = 4, 7, 9, 10, 17, 19 days were examined to determine the time-dependent inhibition.

The inhibitory effect of anthocyanin (>100 mg cyanidin glucoside/100 g plant) was determined using 14 µg and 58 µg extract/38 g chick embryo (Figure 2A). Anthocyanin-treated chick embryo group I (14 µg anthocyanin/chick embryo) showed decreased vessel formation compared to the vehicle-only control when it was administered at pre-incubation embryonic day (ED) = 2 and analyzed at post-injection embryonic day 8 (ED = 10). Higher concentration of anthocyanin-treated group II (58 µg anthocyanin /38 g chick embryo) significantly inhibited blood vessel formation compared to vehicle-only control or lower concentration of anthocyanin group I. Other kinetic time point, including pre-incubation ED = 4 and post-injection ED = 6, was examined (Figure 2B). At ED = 10 chick embryo was harvested and vessel formation was analyzed. Our results demonstrated that increased concentration of anthocyanin rich extract led to significant decrease in the angiogenic index determined through counting the number of blood vessel branch points. We observed maximum inhibition of blood vessels formation at embryonic day ED = 2 compared to treatment at later stage (pre-incubation ED> 10 days). When anthocyanin was administered at later stage with pre-incubation ED = 9 and post-incubation ED = 8, no phenotypical differences were observed (Figure 2C). To determine early time-dependent anti-angiogenic index by anthocyanin extract, we injected anthocyanin at pre-incubation day ED = 0 and post-incubation day ED = 4. Angiogenic index showed 60 % inhibition of vessel length growth compared to control (Figure 3) supporting the hypothesis that early treatment (ED = 0) of anthocyanin on the embryo significantly (p< 0.05) inhibited blood vessel length growth compared to the control.
a quantitative evaluation of the vessel network by extracting characteristic information of the angiogenic images. Parameters analyzed were inhibited by anthocyanin treated chick embryo (anthocyanin + = 14 µg anthocyanin and anthocyanin ++ = 58 µg anthocyanin/38 g chick embryo), compared to the control (Figure 4B). We determined the angiogenic parameters, including extremities, nodes, junctions, master junction, master segments, master segments length, meshes, total meshes area, number of segments, isolated elements and branches, number of segments, branches, number of isolated elements, total length, total branching length, total segments length, total branches length, total isolated branches length, branching interval, total segments length/branches, mesh index, master segments length/master segments and mean mesh size. Next, we examined Angiogenesis Analyzer to obtain angiogenic index quantitatively (Figure 5). Systematic analysis showed extremities (A1-B1), nodes identified as 3 neighbor (A2-B2), twig (C1, D1), segment (C2, D2) delimited by two junctions (C3, D3), and branch (C4, D4). E shows a junction implicated only in branch (E1) and master junctions like E2 delimiting master segments (E3). F represents the master tree composed from master segments associated by master junctions delimiting the meshes (F1). Two close master junctions can be fused into a unique master junction (F2) and the underlying segment (F3).

3.3 Fatty Acids as an Angiogenic Stimulators

Fatty acids that include eicosanoid may serve as effective pro-angiogenic factors, considering that fatty acids regulate vascular endothelial cell proliferation, migration and capillary formation [14]. Anti-angiogenic anthocyanin was compared to selected fatty acids mixtures as angiogenic stimulator (Figure 6). Fatty acids, including oleic (18:1), linoleic (18:2, w6), linolenic (18:3, w3), palmitic (16:0), palmitoleic (16:1), up-regulated vessel length and branches 20-50%, compared to control. Ten major angiogenic indexes, that includes extremities, nodes, junctions, master junction, number of master segments, meshes, total meshes area, number of segments, branches, number of isolated elements, total length, total branching length, total segments length, total branches length, total isolated branches length, branching interval, total segments length/branches, mesh index, master segments length/master segments and mean mesh size. Next, we examined Angiogenesis Analyzer to obtain angiogenic index quantitatively (Figure 5). Systematic analysis showed extremities (A1-B1), nodes identified as 3 neighbor (A2-B2), twig (C1, D1), segment (C2, D2) delimited by two junctions (C3, D3), and branch (C4, D4). E shows a junction implicated only in branch (E1) and master junctions like E2 delimiting master segments (E3). F represents the master tree composed from master segments associated by master junctions delimiting the meshes (F1). Two close master junctions can be fused into a unique master junction (F2) and the underlying segment (F3).
3.4 Molecular Modeling

A 3D model of the protein-ligand interaction was built using the 3D structure of VEGFR2 as a template and selected anthocyanin as a ligand. VEGFR2 was obtained from the Protein Data Bank [20,21]. Six molecules, including cyaniding-3-glucoside and hibiscetin from Hibiscus were selected for protein-ligand modeling analysis (Figure 8a) [22–24].

The conserved residues presenting VEGFR2 active sites were A866, L840, K920, F918, G922, C919, E917, V848, C1045, F1047, L1035, G922, K920, D1046, V916, V899, K868, and E885. The modeling was performed using SwissDock. SwissDock generated many binding modes using EADock software as local docking in the box as well as blind docking in the vicinity of all VEGFR2 cavities. At the same time, binding energies (ΔG) were calculated and the most favorable energies were evaluated to show the most favorable clusters. Total 257 binding were calculated with clusters (0-37), elements (0-15), FullFitness (-1588.73 to -1567.50 kcal/mol), binding AG (-8.42 to -6.20 kcal/mol).

The conserved hydrophobic amino acids in receptor tyrosin kinases including A866, L840, F918, G922, V848, F1047, L1035, G922, V916, and V899 in the active site of VEGFR2, were proposed to interact with three aromatic rings of hibiscetin by forming hydrophobic interaction and ππ interaction (Figure 8b). Other conserved amino acids seem to contribute to forming hydrogen bonding between K920, C919, E917, C1045, K920, D1046, K868, and E885 and 7 hydroxyl groups of hibiscetin. Trihydroxyphenyl ring of hibiscetin interacts with C1045 and D1046 through hydrogen bonding whereas the other two aromatic ring fitted in the hydrophobic pocket of the active site. Hibiscetin contains seven hydrogen bond donor sites and nine hydrogen bond acceptor atoms. Trihydroxyphenyl group is perpendicular to the chromenone structure, thus all three aromatic rings fit into the hydrophobic pocket of VEGFR2.

The molecular mechanism of the proper balance between angiogenic activation and inhibition remains elusive. The current study was designed to investigate whether natural antiangiogenic molecules modulate angiogenesis and how the proper balance of pro- and anti-angiogenic signaling can be analyzed using a chick embryo model [19,25]. Our data demonstrated that systematic administration of anthocyanin-rich extracts of Hibiscus plant blocked collagenolytic activity in ovo, leading to a significant reduction of vessel formation of chick embryo. Systematic analysis of angiogenic index demonstrated that the hierarchical branching pattern of vessel formation exists in the chorioallantoic membrane (CAM) and both global as well as local organization of the vascular network characterizes angiogenesis. Further, the current study introduced the systematic analysis of angiogenic index using the Angiogenesis Analyzer that detects vessel networks and analyzes the vascular organi-
zation of endothelial cells. To evaluate the angiogenic effects of anthocyanin and fatty acids, the Angiogenesis Analyzer obtains a quantitative evaluation of the vessel network by extracting characteristic information of the angiogenic images.

Previously, our proteomic studies under stress conditions identified the following molecular changes in apoptotic and angiogenic reactions in the retina and RPE that include (1) mitochondrial dysfunction in the peripheral RPE in AMD, (2) oxidative stress by intense (>7,000 lux) and constant light (>1 week), (3) cytoskeletal remodeling by actin, vimentin, and tubulin under stress, (4) high concentration of nitric oxide, (5) hypoxia, (6) disrupted circadian clock, (7) erythropoietin changes via pAK2, pSTAT3, Bclxl, HIF1, VEGFR (8) altered lipid concentrations, (9) altered retinoid metabolism (RPE65, CRALBP), (10) altered energy metabolism by S/T/Y kinases, creatine kinase, pyruvate kinase (11) altered expressions of heat shock proteins, crystallins [6,8,9,27]. Based on our proteomic data, we tested the following molecules as anti-apoptotic and anti-angiogenic molecules: (1) prohibitin as an anti-apoptotic mitochondrial-nuclear shuttle, (2) erythropoietin as an anti-apoptotic protein via JAK2/STAT3 pathway, (3) melatonin as an anti-apoptotic, anti-angiogenic molecule, and a circadian determinant protecting cytoskeletal reorganization through PP2A/vimentin pathway, (4) okadaic acid, arginine, and SNAP (nitric oxide generator) to regulate nitric oxide concentration, (5) changes in certain lipids, including cardiolipin and cholesterol as specific biomarkers of apoptotic retina and RPE under stress conditions [10].

The chorioallantoic membrane in a chick embryo model offers a number of unique advantages to study the complex, multi-step process of angiogenesis [19]. The chorioallantoic membrane provides a supportive environment for a source of angiogenic blood vessels. The chorioallantoic membrane comprises the fully developed ectoderm capillary plexus which is a network of tiny capillaries connecting the arterial and venous blood vessel networks. Late injection (t = 10-15 days of pre-incubation) of anthocyanin has less influence than early injection (t = 0 to 4 days pre-incubation). Fatty acids up-regulated vessel length (20-30%) and isolated branches (50%), respectively.

At the later stage, CAM assay represents the certain disadvantage because it may contain a pre-developed vascular network and the vasodilation that invariably follows its manipulation may be hard to distinguish from the effects of the test substance.

**Hibiscus sabdariffa** is identified as an annual red Sorrel dicotyledonous plant. It is an herbaceous tropical plant containing anthocyanin, flavonoids and polyphenol acids which includes cyanidin-3-sambubioside, delphinidin-3-O-glucoside and delphinidin-3-O-sambubioside. Historically *sabdariffa* extract has been used for treating liver disorders, hypertension, pyrexia, diuretic, emollient and purgative. The calyx mixtures decrease intestinal transit, and exert anti-inflammatory and anti-mutagenic effects. Its beneficial effects are linked to the pres-
ence of phytochemical constituents including anthocyanin which is the most abundant (>100 mg cyanidine rutinoside/100 g plants).

**Fig. 8.** A. Two-dimensional and 3-dimensional structures of six different anthocyanins, including delphinidin-3-glucoside, delphinidin-3-sambubioside, cyanidin-3-glucoside, cyanidin-3-samboside, malvidin-3-gallactoside, and hibiscetin, were presented in *Hibiscus sabdariffa* for the molecular binding study. B. A 3D model of the protein-ligand interaction was built using the 3D structure VEGFR2 and hibiscetin as a template. VEGFR2 was obtained from the Protein Data Bank (1Y6A).

SwissDock as a protein-ligand docking server was used for ligand-protein interaction studies (http://www.swissdock.ch/docking). The active site of VEGFR2 and protein structure were examined among seven X-ray diffraction crystal structures (PDB ID = 1Y6A, 1Y6B, 1YW1, 2OH4, 2P2H, 2P21, 2QUS). 257 bindings between VEGFR2 and hibiscetin were calculated with clusters (0-37), elements (0-15), FullFitness (-1588.73 to -1567.50 kcal/mol), binding energy ΔG (-8.42 to -6.20 kcal/mol).

The 10 conserved hydrophobic amino acids, including A866, L840, F918, G922, V848, F1047, L1035, G922, V916, and V899 in the active site of VEGFR2, were proposed to interact with three aromatic rings of hibiscetin by forming hydrophobic interaction and π-π interaction (Figure 8b). Other conserved amino acids seem to contribute to forming hydrogen bonding with K920, C919, E917, C1045, K920, D1046, K868, and E885 and hydroxyl group of hibiscetin. Trihydroxylphenyl ring of hibiscetin interacts with C1045 and D1046 through hydrogen bonding whereas the other two aromatic ring fitted in the hydrophobic pocket of the active site. Hibiscetin contains seven hydrogen bond donor sites and nine hydrogen bond acceptor atoms.

The current study determined 21 angiogenic index, including vessel length, branches, nodes and segments, using angiogenesis analyzer to show 50-60 % inhibition of blood vessels formation by anthocyanin rich extract of *H. sabdariffa*. Our results also revealed that selected fatty acid mixtures that include oleic (18:1), linoleic (18:2, w6), linolenic (18:3, w3), palmitic (16:0), and palmitoleic (16:1) acid, significantly stimulated angiogenesis by 20-50 %. Our data suggest that conjugated double bond-containing natural products from sub-Saharan African plants library may regulate the vessel formation in time- and dose-dependent manner. Our studies imply that potential molecular determinants including anthocyanin may inhibit angiogenesis and the proper balance of pro- and anti-angiogenic signaling may control the vascular microenvironment. The current study demonstrated that anthocyanin rich extract of *H. sabdariffa* has anti-angiogenic effect in vitro and may be a potential candidate for the treatment of angiogenic diseases, including wet AMD.

**4 Conclusion**

The current study examined the anthocyanin-rich Hibiscus extracts to determine their time- and dose-dependent anti-angiogenic properties using chick chorioallantoic membrane
model in ovo. Using this well-characterized model we showed that anthocyanin-rich extracts inhibited angiogenesis in time and concentration-dependent manner showing maximum inhibition of vessel formation in early treatment on 0-2 p re-incubation day. Quantitative determination of angiogenesis using 21 angiogenic indexes provided the evidence that Hibiscus sabdariffa inhibits angiogenesis and therefore can be used for treating unco ntrolled angiogenesis-related diseases, including wet AMD. Imbalance between angiogenic stimulators and inhibitors may lead to incomplete vessel maturation. Our data suggest that angiogenic determinants that include fatty acids and anthocyanin as well as th eir concentration, pre-incubation timing and injection timing may exist in neovascularigenesis and angiogenesis.

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