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The Role of the Estrogen-Related Receptor Alpha (ERRa) in Hypoxia and Its Implications for Cancer Metabolism

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Abstract: Under low oxygen conditions (hypoxia), cells activate survival mechanisms including metabolic changes and angiogenesis, which are regulated by HIF-1. The estrogen-related receptor alpha (ERR α) is a transcription factor with important roles in the regulation of cellular metabolism that is overexpressed in hypoxia, suggesting that it plays a role in cell survival in this condition. This review enumerates and analyses the recent evidence that points to the role of ERR α as a regulator of hypoxic genes, both in cooperation with HIF-1 and through HIF-1- independent mechanisms, in invertebrate and vertebrate models and in physiological and pathological scenarios. ERR α 's functions during hypoxia include two mechanisms: (1) direct ERR α /HIF-1 interaction, which enhances HIF-1's transcriptional activity; and (2) transcriptional activation by the ERR α of genes that are classical HIF-1 targets, such as VEGF or glycolytic enzymes. ERR α is thus gaining recognition for its prominent role in the hypoxia response, both in the presence and absence of HIF-1. In some models, ERR α prepares cells for hypoxia, with important clinical/therapeutical implications.

Keywords: ERR; HIF-independent response to hypoxia; cancer; metabolic adaptation to hypoxia; VEGF; angiogenesis; ischemia; PGC- 1α

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1. Introduction

Oxygen, the final electron acceptor of the mitochondrial respiratory chain for ATP production, is crucial for all aerobic organisms. Despite the complex systems that higher organisms have developed to irrigate every organ and constantly provide all cells with oxygen, a variety of conditions can limit oxygen levels: some pathological (ischemia, tumor development, anemia, lung disease) and some physiological (embryonic development, exercise). Hypoxia is thus defined as a decrease in the oxygen supply to levels insufficient for cellular function [1]. The precise O2 concentration that represents hypoxia varies from tissue to tissue, and likely even between individuals, as different tissues are exposed to different physiological oxygen concentrations (termed "physioxia"), most ranging from 3 to 9% oxygen (23–70 mmHg) [1]. O2 determinations suggest that most mammalian high-energy-demand tissues, such as the brain, muscle, liver, renal cortex and heart, maintain physiological oxygen concentrations between 2.5 and 5.5% [1,2]. Since hypoxia can quickly become life threatening, it triggers a response to modulate blood flow, change energy metabolism, induce angiogenesis and cell differentiation and, ultimately, induce apoptosis [3].

The hypoxia response has a fast component that relies on existing proteins, such as ion channels and already-expressed signaling pathways, with effects such as blood

redistribution, tachypnea/tachycardia, widespread inhibition of protein translation and impaired cell proliferation [4,5]. It also has a well-characterized slower component that induces the expression of about one-thousand specific hypoxia genes once hypoxia is installed [6]. Hypoxia-inducible factors (HIFs) are central transcription factors responsible for protein expression during the slow component of the hypoxia response [7], but other transcription factors also participate. One such example is the estrogen-related receptor alpha (ERR α), a ubiquitously expressed orphan nuclear receptor, abundant in high-energy-demand tissues such as the heart, kidneys and cancer cells [8–11]. This review discusses the role that ERR α plays in the hypoxia response in synergism with HIF-1 and by HIF-independent mechanisms.

2. HIF-1, -2 and -3 Mediate the Hypoxia Response

The HIF-mediated transcriptional response to hypoxia was discovered in the 1990s [7] and received the Nobel Prize in Physiology or Medicine in 2019 [12]. Due to its central importance to physiology and to pathological states such as cancer, HIF has been extensively studied and reviewed [6,12–15]. HIF-1 was the first such factor described and remains the most characterized, but HIF-2 and HIF-3 have been described as well [14,16]. All are heterodimeric basic helix–loop–helix transcription factors consisting of subunits α and β . Each α subunit is O2-regulated, as it is constantly targeted for destruction during normoxia via the Von Hippel Lindau protein and the E2-ligase/ubiquitin proteasome pathway [12–15,17]. In turn, the HIF-1 β subunit (initially known as the aryl hydrocarbon nuclear translocator ARNT, UniProt P27540) is constitutively expressed [12–15] and can heterodimerize with the different oxygen-sensitive α subunits (HIF-1 α , HIF-2 α or HIF-3 α) to create tissue-specific HIF-1, HIF-2 or HIF-3 transcription factors. While HIF-1 α is conserved from Parazoa to vertebrates and expressed in most cells [13,14], HIF-2 α and -3 α are only present in vertebrates and expressed tissue-specifically [12,14,16].

HIF-1 α (HIF1A, UniProt Q16665), the most characterized homolog, is an 826-residue protein that locates to the cytoplasm during normoxia where it is constantly destroyed [12–15,17]. Under hypoxic conditions or in the presence of iron chelating agents, it translocates to the nucleus and dimerizes with HIF-1 β to form the functional HIF-1 complex that activates gene transcription [12–15,17]. HIF-1 α DNA binding activity and stabilization have half maximal responses between 1.5 and 2% O₂ and maximal response at 0.5% O₂, determined in human cultured cells [2]. HIF-1 requires co-activators/transactivators such as CREB binding protein (CBP), p300 [18,19] and others [14], and the complex binds to hypoxia response elements (HREs) with the 5'-RCGTG-3' consensus [14,20].

Around one-thousand genes have been identified as HIF-1 targets [21] and they can be grouped into two functional categories: those that increase oxygen supply to tissues and those that decrease oxygen consumption by tissues [6,15]. In the first category, HIF elicits an increase in oxygen delivery to tissues by triggering erythropoiesis and angiogenesis through the expression of erythropoietin, the hormone that controls red cell production and blood O2-carrying capacity [7], and VEGF (vascular endothelial growth factor), the main protein that stimulates new blood vessel formation [22]. In the second category (decreasing oxygen consumption) are many genes that modify energy metabolism [23,24]. For example, in hypoxia, oxidative phosphorylation (OXPHOS) is restricted due to the lack of O2, and cells shift to anaerobic glycolysis through the increased expression of glycolytic enzymes by HIF-1 [23]. To compensate for the much lower ATP generation per glucose molecule through glycolysis than through OXPHOS, HIF-1 activates transcription of the SLC2A1 and SLC2A3 genes coding the glucose transporters GLUT1 and GLUT3 that increase glucose uptake [23]. Moreover, to inhibit the conversion of pyruvate to acetyl CoA, HIF-1 activates gene transcription to decrease pyruvate flux to the Krebs cycle and increase lactate production. Examples of activated genes are the PDK1 gene encoding PDH kinase, which phosphorylates and inactivates the catalytic subunit of pyruvate dehydrogenase (PDH) [24], and the LDHA gene encoding lactate dehydrogenase A, which directly catalyzes the conversion of pyruvate to lactate [20]. In this way, the HIF-1 response to hypoxia is, in part, executed through metabolic adaptation.

To form HIF-2, HIF1- β heterodimerizes with HIF-2 α (EPAS1, UniProt Q99814; also called endothelial PAS domain protein 1, HIF-1 α -like factor (HLF), HIF-1 α related factor (HRF) and member of the PAS superfamily-1 (MOP-1)). HIF-2 α has similarities to HIF-1 α in terms of domain structure, O2-dependent degradation, DNA sequence recognition (also binds to hypoxia response elements, HREs) and heterodimerization, yet exhibits different effects over gene expression mostly due to tissue-specific expression and kinetics [12,14,16]. The kinetics of HIF-1 α and HIF-2 α suggest that the former exerts a more rapid response at oxygen levels around 1–2%, whereas HIF-2 α action occurs after prolonged hypoxia [25]. In contrast to HIF-1 α that expresses ubiquitously, HIF-2 α only expresses in certain tissues such as embryonic and adult vascular endothelia, lung, placenta, heart, renal interstitial cells and liver [16,26]. HIF-2 α also has specific coactivators such as NF- κ B essential modulator, and Ets1 that do not interact with HIF- 1α . Genes strongly activated by HIF-2 are erythropoietin, VEGF receptor 2, insulin-like growth factor-binding protein-2 and plasminogen activator inhibitor-1 [26,27]. HIF-2 acts more effectively than HIF-1 on erythropoietin and iron metabolism genes, whereas VEGF and GLUT1 are similarly activated by HIF-1 and HIF-2, and glycolytic enzymes are more activated by HIF-1 [26,27]. Thus, within the hypoxia response that requires gene expression, HIF-1 constitutes a faster metabolic component; in turn, HIF-2 is more effective on erythropoiesis control once hypoxia persists.

In turn, HIF-3 α (HIF3A, UniProt Q9Y2N7) can be present in different splice variants that depend upon the tissue, some of which are proposed to have negative regulatory functions on the hypoxia response [16,28]. This seems to be the case for the short HIF-3 α variant that is also called inhibitory PAS domain protein (IPAS), which expresses in corneal epithelium and putatively prevents vascularization there [16,28]. Thus, HIF-3 α could have specific regulatory roles that will not be further reviewed here, but that have been discussed in [16,28].

3. Introduction to the Estrogen-Related Receptors (ERR)

The orchestration of metabolic adaptation central for hypoxia survival seems to involve other transcription factors that specialize in the control of energy metabolism, such as the estrogen-related receptor (ERR) subfamily of nuclear receptors. Here, we describe the subfamily, with a focus on ERR α , and then analyze the evidence and mechanisms that link ERR α to the regulation of hypoxic metabolism and angiogenesis.

The ERR subfamily belongs to group III of the nuclear receptor superfamily (orphan nuclear receptors) [8,29]. In humans and most vertebrates, it comprises three members: ERR α (NR3B1), ERR β (NR3B2) and ERR γ (NR3B3). However, only one ERR gene has been found in invertebrates such as Urochordates, *Drosophila melanogaster*, and in the mosquito *Anopheles gambiae*, but none seem to exist in *Caenorhabditis elegans* [30–32]. A search in the Inparanoid database (version 9) confirmed that no ERR homologs are identifiable in nematodes [33].

The general structure of ERRs is common to nuclear receptors, including four functional domains: N-terminal (NTD), DNA-binding (DBD), hinge, and a putative ligand-binding domain (LBD) [8,10,34] (Figure 1A). The DBD comprises two cysteine-rich zinc finger motifs, which are required for DNA binding and recognize the ERR response elements (ERREs), composed by the sequence TNAAGGTCA [35–37]. The three members of the ERR family (α , β and γ) bear high similarity, particularly in the DBD and LBD domains [38], but they have somewhat different functions and their expression is tissue-specific. ERR γ and ERR β bear more similarity with each other than with ERR α [38]. ERR α is the most abundant member of the family, expressed in most cells, and with higher levels in those with high energy demand, especially in cells that oxidize fatty acids [35–37], compatible with its role in the transcriptional control of energy metabolism.

ERR α (ESRRA, UniProt P11474) is a 423-residue protein and the first orphan nuclear receptor identified in a 1988 screen for genes related to estrogen receptor alpha (ER α) [39], just a few years before the identification of HIF. Unlike estrogen receptors, no endogenous ligand has been described for ERRs; thus, ERR α , the first orphan nuclear receptor identified, remains among the "non-adopted" orphans [40,41]. Recently, it was reported that an endogenous 19-nor steroid estradienolone, found in the urine of pregnant women, can bind and act as an inverse agonist to ERR α and ERR γ [42]. There is still little information to discern if this could be the long-sought endogenous ligand of the family, but it seems unlikely due to the plethora of crucial functions that have been described for the ERRs and which do not require a ligand (reviewed below).

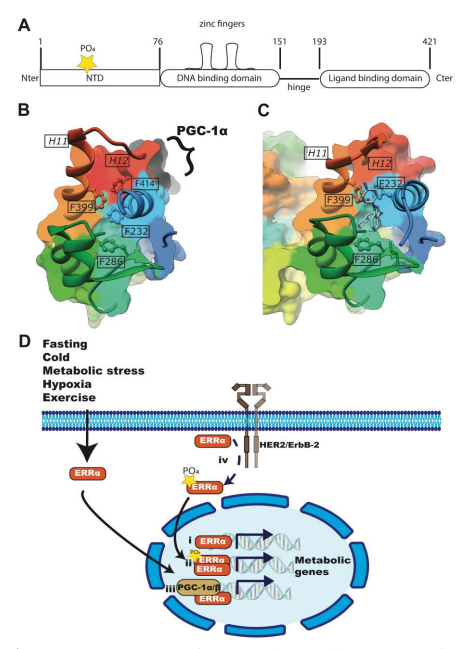


Figure 1. A. ERRα's structure and function. (**A**) The general domain topology of ERRα and other estrogen-related receptors (NR3) includes an amino-terminal domain (NTD), a DNA binding domain, a hinge, and a putative ligand-binding domain (LBD). (**B**,**C**) ERRα's LBD has been crystalized in the absence (**B**) and presence of ligand. (**C**) The binding site is lined by residues F232, F286, F399 and F414 (PDBIDs 1XB7 and 2PJL) providing bulky side chains that fill the ligand-binding pocket in the absence of ligand (**B**). Moreover, without ligand, helix 11 (H11) lifts away from the binding site; helix 12 (H12) is perpendicular to H11 and residue F414 occludes the binding site (**B**). (**D**) ERRα can

bind DNA as a (i) monomer or (ii) dimer. ERR α 's transcriptional activity increases in complex with co-activators, for example PGC-1 α or β (iii), and through posttranslational modifications, such as phosphorylation (iv), mediated by HER2-EGF. ERR α activity increases in metabolic stress, cold, fasting/nutrient deprivation, exercise and hypoxia.

Due to ERR α 's sequence identity to ER α , particularly in the DBD and LBD (68% and 37% residue identity, respectively [39]), it was initially suggested that these two receptors share common targets, co-regulatory proteins and sites of action [43,44]. However, through the combination of computational biology; ERR α silencing; interaction with the co-activators such as PGC-1 α ; DNA binding assays; chromatin immunoprecipitation with sequencing (CHIP-seq); and reporter gene approaches, the differences between ERR α 's and ER α 's functions have become apparent [8,36,37,45]. ERR α regulates a different set of genes to Er α and is not involved in estrogen response. ERR α is mainly involved in the transcriptional regulation of metabolic pathways spanning carbohydrate, lipid and amino acid metabolism, importantly through the regulation of genes for mitochondrial biogenesis, oxidative phosphorylation (OXPHOS) and fatty acid oxidation [8,10,11,37]. The other members of the ERR family also control aspects of metabolism, although in specific tissues [8,29,32,38]. Overall, ERRs occupy the promoters of over 700 genes that encode mitochondrial proteins, regulating mitochondrial biogenesis [8].

Specifically, ERR α binds to the promoters of glycolysis and tricarboxylic acid cycle (TCA) genes, and to OXPHOS genes such as ATP synthase b (ATPsynb), cytochrome c (CYCS), COX4, GABPA and adenine nucleotide translocator 1 (ANT1) [45]. ChIP-seq studies performed in mouse or human, liver, kidney, macrophages or cancer cell lines confirmed that ERR α can bind to promoters for OXPHOS (*Sdhd and Sucla2*), TCA (*Fasn*), glycolysis/gluconeogenesis and lipid metabolism genes (*Gpam and Elovl6*) [37,46–49]. Furthermore, ERR α activates the promoters of β -oxidation genes, such as ACADM (mediumchain acyl co-A dehydrogenase) and CPT1A (carnitine palmytoyl transferase 1A), as well as the promoters of glutamine transporters and enzymes for glutamine synthesis and catabolism [35,45]. In summary, ERR α 's function can be described as activating gene expression to adapt energy production to physiological or pathological stress. ERR α 's functions in cellular metabolism have been reviewed in [8]. In breast cancer, ERR α 's transcriptional activities mediate metabolic adaptations leading to treatment resistance [47]. The metabolic programs it controls make ERR α an ideal contributor to the hypoxia response and a potential pharmacological target.

The other members of the ERR subfamily also modulate metabolism with complementary and sometimes opposite functions to ERR α [8,29]. ERR β has emerged as important in maintaining multipotency [38]. In breast cancer, ERR α and ERR γ seem to play opposing roles as modulators of cell metabolism: ERR γ activates TCA and OXPHOS while ERR α redirects energy metabolism to glycolysis and lactate production [8]. This balance of control is likely part of the mechanism at the core of the Warburg effect in many tumors, along with HIF-1 [8,11], but it is far from a simple on/off switch. Rather, it is a dynamic balance under tight control that is highly cell- and context-specific, where both members of the ERR family, ERR α and ERR γ , activate metabolic pathways facilitating cell survival and adaptation to the changing environment.

To bind DNA and to modulate target genes, ERRs can act as monomers, homodimers or heterodimers [36,37] (Figure 1D), although in live cells mainly homo or heterodimers have been associated to function [37,38]. ERR transcriptional activity is increased by members of the steroid receptor co-activator (SRC) family [43,50,51] and by the peroxisome proliferator-activated receptor gamma co-activator-1 (PGC-1) α and β [45,50] (Figure 1D). Interactions with the cofactors are mediated by ERR's LBD, particularly by helices 11 and 12 via leucine rich motifs (H11 and H12 in Figure 1B,C), also referred to in the literature as ERR's AF2 domain for "activation function 2" [50,53,54]. In particular, ERR α 's functions on metabolism are more dependent on PGC-1 α [50]. ERR α and PGC-1 α influence each other's expression, and both orchestrate the transcription of energy metabolism genes

[37,54]. Recently, details on ERR α 's transcription initiation mechanisms have been clarified. PGC1 α was essential for p300 and mediator recruitment to activate transcription when ERR α acted on chromatin, whereas on naked DNA ERR α established direct contact with initiation factor TFIIH, and PGC1 α did not further increase transcription [50,51]. While ERR α depends on PGC-1 α to transcribe metabolic genes, ERR β and γ can function independently of PGC-1 α in stem cells and muscle [50], and other cofactors that interact with their AF2 domains, such as NCOA, replace PGC-1 α [50,51].

Unlike ER α , the ERRs do not need ligands to interact with its co-activators and to bind DNA (they are constitutively active), probably because the putative ligand-binding pocket (LBP) is occupied by residue side chains in a conformation favored by binding to the cofactors. In the empty ERR α crystal structures, the binding pocket is mainly occupied by the bulky phenolic ring of Phe232 (XRD structures number this residue as 328; however, numbering according to UniProt is used here), which corresponds to a less bulky Ala350 in ER α [41]. Despite this apparent lack of a prominent ligand-binding pocket [41,53], synthetic compounds can inhibit ERR α 's constitutive activity; thus, they are considered ERR α 's inverse agonists (i.e., compounds with affinity and intrinsic activity on the protein). Among the first synthetic inverse agonists described for ERR α was XCT790 (reported in 2004), a thiadiazole acrylamide, which alters ERR α /PGC-1 α signaling and is inactive against the rest of the ERRs and ER α [55]. Later, "compound 1a" (ciclohexilmetil-(1-p-tolil-1H-indol-3-ilmetil)-amine) and "compound 29" (4-(4-{[(5R)-2,4-dioxo-1,3-thiazolidin-5-yl]methyl}-2-methoxyphenoxy)-3-(trifluoromethyl)benzonitrile) were synthetized and have been crystallized in complex with ERR α 's LBD [56,57]. An analysis of these ERR α structures with inhibitors revealed a significantly larger ligand-binding pocket than in the empty LBD, created by the rearrangement of amino acid residues F232, F286, F399 and F414 (328, 382, 495 and 510 in XRD 2PJL and 1XB7). F232 and F414 change conformation significantly when ERR α admits a ligand (Figure 1B vs. Figure 1C). In addition, these structures suggest that the presence of the inverse agonists disrupts the interaction between ERR α and PGC-1 α , through the displacement of ERR α 's helix, to a position that interferes with co-activator recruitment [56,57].

4. Evidence of ERR α 's Participation in the Hypoxia Response

Next, we review the evidence for $ERR\alpha's$ participation in the hypoxia response in models that span invertebrates and vertebrates, and physiological and pathological scenarios. These studies have led to the discovery of HIF-dependent and independent mechanisms, including some that transcend $ERR\alpha's$ central role as a metabolic coordinator during stress.

4.1. ERRα Induces VEGF Expression during Muscle Ischemia and Other Models

The work that first pointed to ERR α 's role in hypoxia came from the study of angiogenesis/ischemia where VEGF, a classical HIF-1 target central to angiogenesis, was discovered to also be inducible by ERR α in skeletal muscle [58]. Arany et al. first detected that ERR α 's co-activator, PGC-1 α , was induced by hypoxia in vitro in various cell types, and in vivo in muscle [58]. Using a skeletal muscle ischemia model, these authors showed that transgenic animals overexpressing PGC-1 α had increased angiogenesis with VEGF expression. PGC-1 α /ERR α , but not other transcription factors co-activated by PGC-1 α , were necessary to increase VEGF expression, through a mechanism that neither depended on HIF response elements (HREs) nor affected HIF-1 expression/stability [58]. Furthermore, conserved ERR α response elements (ERRES) were identified in the first intron of the VEGF gene and were recognized by PGC-1 α /ERR α [58]. ERR α 's ability to induce VEGF expression and angiogenesis, as well as platelet-derived growth factor (PDGF) and Angiopoietin 2, has been confirmed in other models [59–64]. Some studies suggest that the effect does not require HIF-1 [58,59], while others suggest that HIF-1 can increase ERR α expression [61], and that, in turn, ERR α suppression can decrease HIF-1 α [60].

In skeletal muscle, the alternatively spliced truncated isoforms of PGC-1 α , NT-PGC-1 α and PGC-1 α 4, induced VEGF expression by ERR α without increasing mitochondrial biogenesis [64] (Figure 2B). These PGC1 α isoforms bind ERR α but not other transcription factors, such as NRF-1 and NRF-2 [53,64]. This suggests a mechanism by which the PGC-1 α /ERR α axis can operate in hypoxia without increasing mitochondria (Figure 2A), which would likely be impaired in respiration due to the limited O₂ to act as a terminal acceptor for OXPHOS. Additionally, other authors have suggested that PGC-1 α could amplify intracellular hypoxia by activating mitochondrial biogenesis/OXPHOS as a mechanism to consume all remaining intracellular oxygen [65], thus precipitating hypoxia responses and stabilizing HIF-1 α .

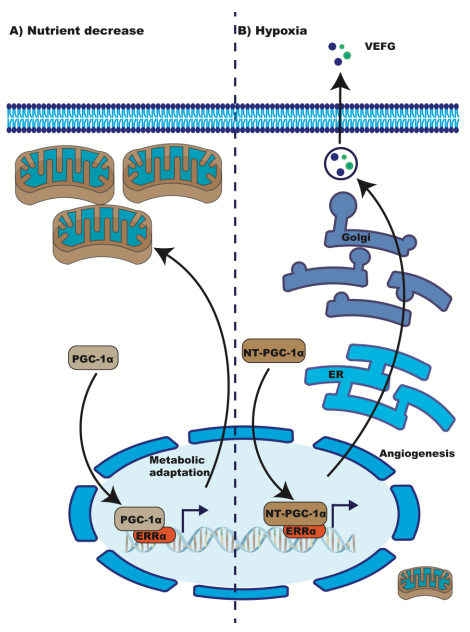


Figure 2. Metabolic adaptation via $ERR\alpha/PGC1\alpha$ in normoxia (**A**) vs. hypoxia (**B**). (**A**) In normoxic nutrient deprivation, $ERR\alpha/PGC1\alpha$ activate mitochondrial biogenesis. (**B**) In hypoxia, $PGC-1\alpha$'s truncated isoform NT-PGC-1 α binds $ERR\alpha$ but prevents the engagement of other transcription factors, limiting mitochondrial biogenesis and favoring the expression of angiogenesis genes such as VEGF. Oxygen consumption by mitochondria favors local hypoxia with concomitant HIF-1 α stabilization.

Other recent studies suggest that some ERR α 's responses to hypoxia in the skeletal muscle are dependent on HIF-1. ERR α is expressed during hindlimb muscle ischemia. Transgenic mice overexpressing ERR α in the skeletal muscle have faster revascularization with more muscle capillaries and higher artery/arteriole density after ischemia [66,67]. ERR α overexpression was also induced in C2C12 myotubes by oxygen deprivation (culture in 95% nitrogen, 5% CO₂), hypoxia-mimetics such as dimethyl-oxaloylglycine (DMOG) or cobalt chloride (CoCl₂), or by nutrient deprivation [66]. Further in vitro experiments showed that ERR α regulates angiogenic gene expression through promoter recognition in C2C12 myotubes, and pointed out that ERR α 's expression was HIF-1-dependent [66], for which the authors predicted 12 putative HIF1A::ARNT response elements in the ERR α gene promoter [66]. Altogether, these authors suggest that HIF is involved in the hypoxic induction of ERR α in the skeletal muscle through the transcriptional regulation of ERR α expression. However, ERR α 's activity was not explored under HIF-1 depleting or activating conditions.

4.2. ERRα in Brain and Spinal Cord Hypoxia/Ischemia

Studies with astrocytes treated with CORM2, a CO-releasing compound that imitates ischemic brain injury, showed that $ERR\alpha/PGG1\alpha$ can increase VEGF expression independent of HIF-1 (that is, even in HIF-1 α -deficient cells) [68]. The treatment induces Heme Oxygenase-1 (HO-1) expression and its metabolites (CO and bilirubin) and promotes Ca²⁺ influx through L-type Ca²⁺ channels producing CaMKK β -mediated AMPK α activation [68]. AMPK α increases NAMPT expression and NAD+ synthesis, which in turn increases SIRT activity. PGC-1 α can be deacetylated by SIRT1 [68], and once deacetylated it interacts with ERR α to increase mitochondrial biogenesis and oxygen consumption [68]. With this model, the authors previously suggested that oxygen consumption aggravates intracellular hypoxia, allowing HIF-1 α stabilization that further increases ERR α /PGG1 α expression [61]. Using ChIP assays, the authors proposed that HIF-1 can stimulate ERR α 's transcription by binding to a putative HIF-1 response elements (+539 to +542, 5'-CGTG-3') within the promoter region of the ERR α gene [61]. HIF-1 α knockdown blocked ERR α 's expression but not PGG1 α 's in that HO-1 inducing model. Therefore, it is likely that HO-1 can stimulate VEGF both via HIF1-α dependent and independent mechanisms, the latter involving PGC-1α/ERRα and calcium regulation through the Ca²⁺/CaMKK/AMPK pathway [68]. These authors also propose some reciprocal and dynamic coordination between HIF- 1α , and PGC- 1α /ERR α for VEGF expression in astrocytic ischemia, involving mitochondrial biogenesis [61].

Other processes such as spinal cord injury (SCI) can manifest with ischemia, which aggravates secondary injury and neurological dysfunction [69–71]. Therefore, the vascular response is critical for SCI repair and includes HIF-1 α and VEGF expression. In an SCI rat model, ERR α inhibition with XCT790 decreased VEGF and angiopoietin-2 expression [72], which in turn decreased endothelial cell proliferation, vascular density and produced histopathological changes to the spinal cord, such as inflammatory cell infiltration, hemorrhage and vacuolation, and fewer normal neurons, suggesting that ERR α activity is essential for SCI repair, in part by favoring adequate re-vascularization via VEGF [72]. In this model, it has not been explored whether ERR α 's effects require HIF-1.

In the microglial cell line, BV2, pharmacological ERR α inhibition (with XCT790) or activation (with pyrido [1,2- α]-pyrimidin-4-one) were explored in combination with CoCl₂ to mimic the hypoxia that accompanies SCI. Hypoxia induced HIF-1 α and autophagy. ERR α 's effects were similar with/without hypoxia although more pronounced in hypoxia. During hypoxia, ERR α inhibition increased autophagy markers and increased IL-6, TNF- α and IL-10 mRNAs, but decreased FNDC5 (fibronectin type III domain containing protein 5) expression. In turn, ERR α 's activation decreased p38 MAPK phosphorylation. The authors suggest that ERR α helps maintain homeostasis in microglia during hypoxia by down-modulating autophagy and inflammation [73].

4.3. ERRα in Hypobaric Hypoxia

On the other hand, in a non-pathological process such as exposure to high altitude, the expression of ERR α and PGC-1 α are downregulated and the cell suffers mitochondrial dysfunction [74]. Treatment with dexamethasone maintains ERR α and PGC-1 α levels similar to normoxia. This effect allows adaptability to hypobaric hypoxia through the expression of ERR α transcripts, Fis1, Drp1 and Mfn2, which are mitochondrial dynamics proteins, and in turn increases OXPHOS [74]. This suggests that ERR α -mediated protection of mitochondrial bioenergetics is required for adaptation to hypobaric hypoxia.

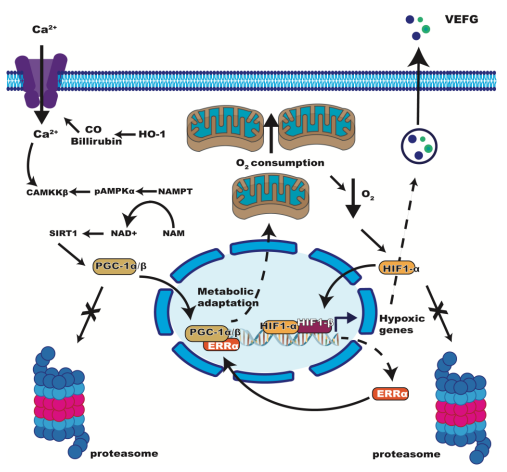


Figure 3. Proposed interactions between HIF-1 and ERR α pathways in hypoxia. ERR α / PGC-1 α express hypoxia genes acting with HIF-1 or independent from HIF-1. Ca²⁺ activates CaMKK β and AMPK α , increasing SIRT. PGC-1 α is activated by SIRT1 deacetylation, decreasing its destruction through the proteasome.

4.4. ERRα's Role in Cancer-Related Hypoxia

In parallel, ERR α has been extensively studied in solid tumors where blood vessels frequently become limiting to irrigate the tumor mass, leading to hypoxia. Thus, cancer represents another model where extensive evidence points to ERR α 's contribution to the hypoxia response. Cancer cells in solid tumors use typical HIF-1 orchestrated mechanisms to survive hypoxia [75], impacting angiogenesis, cancer stem cell maintenance, metabolic reprogramming, epithelial–mesenchymal transition (EMT), invasion, metastasis and resistance to therapy (radiation and chemotherapy) [75–77].

Simultaneously, the overexpression of ERR α has been associated with tumor aggressiveness and poor prognosis [78–80]. In 2002, Ariazi et al. suggested ERR α as a biomarker of unfavorable clinical prognosis in breast cancer, due to increased ERR α mRNA levels in primary tumor cells against normal mammary epithelial cells. ERR α 's expression correlated with Her2/ErbB2, a tyrosine kinase receptor amplified in 15% to 25% of breast

cancers that also confers aggressiveness [78] and that increases ERR α 's transcriptional activity via phosphorylation through MEK/MAPK and PI3K/Akt [80,81].

Subsequent immunohistochemical analyses, mRNA quantification and gene expression profiles in several solid tumors (breast, cervix, colon, endometrium, ovary and prostate) are in agreement with Ariazi et al. and relate ERR α overexpression to cancer aggressiveness, increased risk of recurrence and lower survival [9,77,82–86].

In breast and prostate cancer, ERR α has been found to interact directly with HIF-1 with two main effects: (1) HIF-1 stabilization; and (2) an increase in the HIF-dependent expression of hypoxic genes [86,87]. This evidence has led to the suggestion that the direct ERR α -HIF interaction is another important mechanism by which ERR α contributes to the hypoxic response. The physical interaction between HIF and ERR α has been explored using a series of ERR α truncation mutants covering the N terminus, DBD, and LBD in GST pull down assays. Ao et al. suggested that the ERR α 's DBD is involved in HIF binding [87]. The immunoprecipitation of cellular lysates with anti-ERR α , from MDA-MB-435 breast cancer cells treated with the iron chelator dipyridyl (DP) to stabilize endogenous HIF- 1α , showed that all three ERRs associate to HIF α/β heterodimers both in vitro and in vivo, and this was abolished in ERR α mutants or with ERR α inhibitors [87]. Subsequent studies using co-immunoprecipitation and FRET in prostate cancer cells confirmed that the interaction happens and that it increases HIF-1's transcription but disagree on the ERR α domains involved [86]. Zou et al. suggest that the domain required for interaction with HIF-1 is the AF-2 region in ERR α 's LBD [86]. These authors further suggest that ERR α 's co-activator PGC1 α may be necessary for its interaction with HIF-1, as the AF-2 region in ERR α 's LBD is important for co-activator recruitment. These effects are prevented in ERR α knockdowns or with ERR α 's inverse agonist XCT790 [86,87]. Evidence in prostate cancer cells suggests that the ERR α /HIF-1 α interaction reduces the proteosomal degradation of HIF-1 α [86]. These authors suggest that ERR α overexpression stabilizes HIF- 1α and enhances HIF-1 transcriptional activity even under normoxia, with these effects amplified in hypoxia, resulting in a mechanism for the pre-adaptation to hypoxia [86].

In parallel, Stein et al. reported that ERR α regulates VEGF expression in breast cancer cell lines [62], similar to what was described in the previous section in angiogenesis models. A modified PGC1 α that only binds to ERR α was used to induce VEGF expression in MDA-MB231 and MCF7 breast cancer cells, and this effect was abolished with ERR α knockdowns. A main ERRE was located within the transcribed region of the VEGF gene [62]. The positive regulation of VEGF by ERR α has also been observed in human breast tumors and in murine models [63], supporting that VEGF is a direct transcriptional target of ERR α in cancer, as in other cell types (reviewed in previous section).

In summary, ERR α overexpression enhances the hypoxia response in solid tumor models. It is likely that ERR α functions as an aggressiveness factor in cancer because it prepares cancer cells to resist metabolic stress and hypoxia. ERR α has been observed as active in immunosuppressive and immunoresistant tumors [88]. Cancer models have pointed to HIF-dependent mechanisms such as the physical interaction between ERR α /HIF-1 α , as well as HIF-independent mechanisms such as VEGF modulation by ERR α .

4.5. ERRα and Kidney Hypoxia

Organs with high energy demand such as the brain, heart and kidney have low tolerance to hypoxia and are good models to evaluate ERR α 's effects. Physiological oxygen gradients across the renal cortex and medulla participate in the mechanisms to concentrate urine [89,90]. The healthy human kidney cortex presents around 50 mmHg of oxygen pressure, while the medulla has much lower oxygen pressures between 10 and 20 mmHg [89]. Keppner et al. recently evaluated the transcriptome during hypoxia (24 h at 0.2% O₂) of the cortical kidney murine cell line mCCD(cl1) [90]. They found over 3000 differentially expressed genes, many related to aerobic metabolism and ATP production through

mitochondria, and the hypoxia response was mainly driven by HIF-1 and not HIF-2. Interestingly, they knocked down ERR α and identified a reduced expression of some genes that typically function in hypoxia, such as Egln3 (an alpha-ketoglutarate dependent hydroxylase that controls cell proliferation and transcription upon hypoxia) and Serpin1 (plasminogen activator inhibitor-1, involved in the control of blood clotting) [90]. Since this regulation happened without a change in HIF-1 α , the model suggests that ERR α controls the expression of specific genes important for the hypoxia response.

4.6. Hypoxia in the Invertebrate Fly Model

 $D.\ melanogaster$ is tolerant to oxygen starvation and can survive hypoxia for long periods of time. As in humans, the hypoxia response is importantly mediated by HIF (called sima in $D.\ melanogaster$); thus, $D.\ melanogaster$ has been a study model for hypoxia [91] and represents a vertebrate model with a recognizable ERR. Li et al. showed that, in addition to HIF, the single ERR present in flies (called dERR) is necessary for the hypoxic response in $D.\ melanogaster$, since less than 25% of dERR mutant flies survived hypoxia [92]. Using single and double dERR and dHIF-1 α mutants, they described genes sets that are important for hypoxia response and detected a subset of 282 dERR-dependent transcripts that are HIF-independent and whose expression changed in hypoxia, such as Pgi, Pfk, GAPDH2 and LDH [91]. This work suggests that the dERR has a prominent HIF-independent role in hypoxia adaptation, particularly via the upregulation of glycolytic enzymes.

Additionally, dERR was found to bind dHIF and participate in the HIF-mediated expression of its subset of genes [92]. The binding was shown by two hybrid screen and GST pull-downs and required dHIF's residues 1289–1293 (LKNLL) and dERR's LBD [92], in accordance with Zou et al. in cancer cells [86].

5. Conclusions

HIF-1 is the transcription factor usually considered the main regulator of the hypoxia response. However, ERR α , a cellular metabolism regulator, also plays a key role in hypoxia survival, in models ranging from invertebrates to vertebrates and in physiological and pathological scenarios. ERR α 's functions in hypoxia in most models include two mechanisms: (1) direct ERR α /HIF-1 interaction, which enhances HIF-1's transcriptional activity at HREs (possibly without ERR α 's direct interaction with DNA); and (2) transcriptional activation by the ERR α of genes that are classical HIF-1 targets, such as VEGF or glycolytic enzymes. The second mechanism can even happen in a HIF-1 independent manner that depends on ERREs coexisting with HREs.

ERR α is thus gaining recognition for its prominent role in the hypoxia response, both in the presence and absence of HIF-1. In many models, ERR α prepares cells for hypoxia, with important clinical/therapeutical implications and perspectives that could allow for the manipulation of tissues so they are pre-adapted to resist hypoxia. This is important, as hypoxia is central to numerous diseases with significant human mortality and high costs, such as cancer, cardiovascular and pulmonary disease, stroke, bacterial infections, inflammation, disorders related to prematurity and wound healing.

ERR's expression and activity are conserved from flies (and likely nematodes) to mammals, suggesting that the ERR-mediated response to hypoxia appeared early in evolution. Phylogenetic exploration of the ERR-HIF interaction warrants more interrogation, with the potential to yield insights into its mechanisms and how they evolved.

Despite the many models that have described ERR α as responding to ischemia, it is unknown if ERR α may directly sense oxygen. No mechanism for direct oxygen sensing by ERR α has been described. Alternatively, ERR α 's activation upon ischemia/hypoxia may arise from the metabolic signals derived from ischemia or through HIF-1 stimulation. Protein tyrosine phosphatase 1B (PTP1B) and Parkin have also been shown to, respectively, decrease and increase the transcriptional activity of ERR α in hypoxia models

(pancreatic islets [93] and HeLa cells [94]); thus, other pathways and layers for ERR α modulation likely exist.

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