

Estrogen-related receptor gamma

Estrogen-related receptor gamma (ERR-gamma), also known as [NR3B3](#) (nuclear receptor subfamily 3, group B, member 3), is a nuclear receptor that in humans is encoded by the [ESRRG](#) (EStrogen Related Receptor Gamma) gene.

It behaves as a constitutive activator of transcription.

This protein is a member of the nuclear hormone receptor family of steroid hormone receptors. No physiological activating ligand is known for this orphan receptor, but 4-hydroxytamoxifen and diethylstilbestrol act as inverse agonists and deactivate ESRRG.

It also seems to be the target of bisphenol A.

[Estrogen-related receptors](#) (ERRs) were shown to play an important role in the regulation of [free radical](#)-mediated pathology. This study aimed to investigate the neuroprotective effect of ERRγ activation against early [brain injury](#) (EBI) after [subarachnoid hemorrhage](#) (SAH) and the potential underlying mechanisms. In a [rat model](#) of SAH, the time course of ERRs and [SIRT3](#) and the effects of ERRγ activation were investigated. ERRγ agonist [DY131](#), selective inhibitor [GSK5182](#), or SIRT3 selective inhibitor [3-TYP](#) were administered intracerebroventricularly (icv) in the rat model of SAH. The use of 3-TYP was for validating SIRT3 as the downstream signaling of ERRγ activation. Post-SAH assessments included SAH grade, neurological score, [Western blot](#), [Nissl staining](#), and immunofluorescence staining in rats. In an vitro study, the ERRγ agonist DY131 and ERRγ siRNA were administered to primary cortical neurons stimulated by Hb, after which [cell viability](#) and neuronal deaths were accessed. Lastly, the brain ERRγ levels and neuronal death were accessed in SAH patients. They found that brain ERRγ expressions were significantly increased, but the expression of SIRT3 dramatically decreased after SAH in rats. In the brains of SAH rats, ERRγ was expressed primarily in [neurons](#), [astrocytes](#), and [microglia](#). The activation of ERRγ with DY131 significantly improved the short-term and long-term neurological deficits, accompanied by reductions in [oxidative stress](#) and neuronal apoptosis at 24 h after SAH in rats. DY131 treatment significantly increased the expressions of PGC-1α, SIRT3, and [Bcl-2](#) while downregulating the expressions of 4-HNE and [Bax](#). ERRγ antagonist GSK5182 and SIRT3 inhibitor 3-TYP abolished the neuroprotective effects of ERRγ activation in the SAH rats. An in vitro study showed that Hb stimulation significantly increased intracellular oxidative stress in primary cortical neurons, and DY131 reduced such elevations. Primary cortical neurons transfected with the ERRγ siRNA exhibited notable apoptosis and abolished the protective effect of DY131. The examination of SAH patients' brain samples revealed increases in ERRγ expressions and neuronal apoptosis marker CC3. We concluded that ERRγ activation with DY131 ameliorated oxidative stress and neuronal apoptosis after the experimental SAH. The effects were, at least in part, through the ERRγ/PGC-1α/SIRT3 signaling pathway. ERRγ may serve as a novel therapeutic target to ameliorate EBI after SAH ¹⁾.

¹⁾

Guo Y, Hu Y, Huang Y, Huang L, Kanamaru H, Takemoto Y, Li H, Li D, Gu J, Zhang JH. Role of Estrogen-Related Receptor γ and PGC-1α/SIRT3 Pathway in Early Brain Injury After Subarachnoid Hemorrhage. *Neurotherapeutics*. 2022 Dec 8. doi: 10.1007/s13311-022-01330-8. Epub ahead of print. Erratum in: *Neurotherapeutics*. 2023 Jan 13;; PMID: 36481985.

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