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Novel PPAR- γ agonists as potential neuroprotective agents against Alzheimer's disease: rational design, synthesis, *in silico* evaluation, PPAR- γ binding assay and transactivation and expression studies[†]

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Alzheimer's disease (AD) is a neurological disorder. It is caused by accumulation of amyloid beta (A β) plaques and tau tangles, which gradually leads to cognitive decline and memory loss. Peroxisome proliferator-activated receptor gamma (PPAR- γ), a nuclear receptor, plays a significant role in regulating genes responsible for metabolism and inflammation. Studies have shown that PPAR- γ activation has neuroprotective effects, can potentially reduce inflammation and oxidative stress, and stimulates mitochondrial biogenesis. Current study presents the design, synthesis and *in vitro* evaluation of PPAR- γ agonists for AD that are tailored to optimize binding with the PPAR- γ receptor. The compounds **4a**, **4h** and **4j** exhibited notable binding affinities towards PPAR- γ LBD, with IC₅₀ values of 8.607, 9.242, and 5.974 μ M, respectively, in TR-FRET binding assay. These compounds were cell proliferative and non-cytotoxic in a neuroblastoma cell line (SH-SY5Y). They also demonstrated dose-dependent PPAR- γ activation in transactivation assay. Their neuroprotective effect was studied based on their anti-inflammatory and anti-oxidant potential by reducing the levels of proinflammatory markers (TNF- α , IL-6 and IL-1 β) and ROS in A β -induced SH-SY5Y neuroblastoma cells using a flow cytometry method. The synthesized compounds also showed interactions in molecular docking study with the PPAR- γ receptor and demonstrated good stability in MD simulation. Our results highlight that through activation of PPAR- γ , the compounds **4a**, **4h** and **4j** offer neuroprotective effects by reducing neuroinflammation and oxidative stress, and hence, they may be considered lead molecules for treating AD.

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1. Introduction

AD is a progressive neurodegenerative disorder that largely affects cerebral functions, particularly, memory, thinking and behavior.¹ Neuroinflammation is an inflammation in the brain that contributes to the damage to nerve cells in AD. This inflammation may be triggered by abnormal proteins and immune responses,² and it is characterized by a rapid activation

of microglia, leading to the production of pro-inflammatory cytokines, oxidative radicals, and the infiltration of peripheral immune cells into brain tissue, ultimately contributing to neurodegeneration.³ Aspects such as genetics, chronic diseases and a compromised blood-brain barrier worsen this inflammation. Neuroinflammation accelerates the progression of AD by harming brain cells and promoting the formation of harmful plaques and tangles.⁴ In AD, oxidative stress is linked to A β plaque formation, AD pathophysiological events and synaptic dysfunction. Elevated ROS generation and downregulation of antioxidant defence mechanisms may lead to oxidative stress-mediated neurodegeneration.⁵ Targeting inflammation and oxidative stress through therapeutic interventions may hold promise for the treatment or prevention of AD, although more research is needed to understand the underlying mechanisms and develop effective therapies fully.

PPAR- γ plays a critical role in regulating genes responsible for numerous physiological processes, such as fat cell differentiation, glucose homeostasis, lipid metabolism, and ligand-dependent signal transduction.⁶ PPAR- γ activation is

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associated with anti-inflammatory, anti-oxidant and neuro-protective effects. It improves cognitive function, reduces the accumulation of A β plaques and inhibits the production of inflammatory cytokines, such as interleukin-6 (IL-6) and TNF-alpha (TNF- α), which are elevated in AD that contribute to neuroinflammation.⁷ A β accumulation in the brain activates an immune cell called microglia. PPAR- γ agonists can inhibit the activation of microglia that are induced by A β , thereby reducing neuroinflammation.⁸ They also modulate the processing of amyloid precursor protein (APP), the precursor of A β , and alter the activity of enzymes involved in A β synthesis.⁹ PPARs have been implicated in various neuroprotective mechanisms, including the activation of genes responsible for defending against oxidative stress, promoting anti-inflammatory responses, and stimulating mitochondrial biogenesis.^{10,11}

PPAR- γ activation has advanced as a potential therapeutic avenue in the realm of chronic neurological diseases, such as Alzheimer's disease (AD), multiple sclerosis (MS), amyotrophic lateral sclerosis (ALS), and Parkinson's disease (PD). This is achieved through the modulation of multiple signaling pathways associated with detrimental events in these conditions.¹²

Natural compounds that can activate PPAR- γ include eicosanoids, natural lipid ligands, long-chain fatty acids, nitro-alkenes, and prostaglandins.¹³ However, synthetic ligands called glitazones have gathered significant attention also due to their ability to activate PPAR- γ .¹⁴ Glitazones possess three essential structural features in their binding portion: an acidic head, an aromatic trunk with two carbon linkers, and a lipophilic tail. The ligand binding domain (LBD) of PPAR- γ , which refers to the region of the receptor where ligands attach, is larger than that of most receptors and features a Y-shaped binding pocket that can accommodate diverse ligand structures.¹⁵ When an agonist binds to PPARs, these receptors pair up with RXR receptors to form heterodimers. These PPAR-RXR heterodimers then interact with specific proteins called PPAR-cofactors, such as PGC-1 α , to regulate gene expression.^{16,17} When PGC-1 α is activated by PPAR- γ agonist binding, it enhances gene expression related to energy metabolism, mitochondrial biogenesis, and oxidative stress defense.^{18,19} This leads to improved energy production, reduced oxidative damage, increased fatty acid oxidation, and anti-inflammatory effects. Overall, PGC-1 α activation supports better metabolic

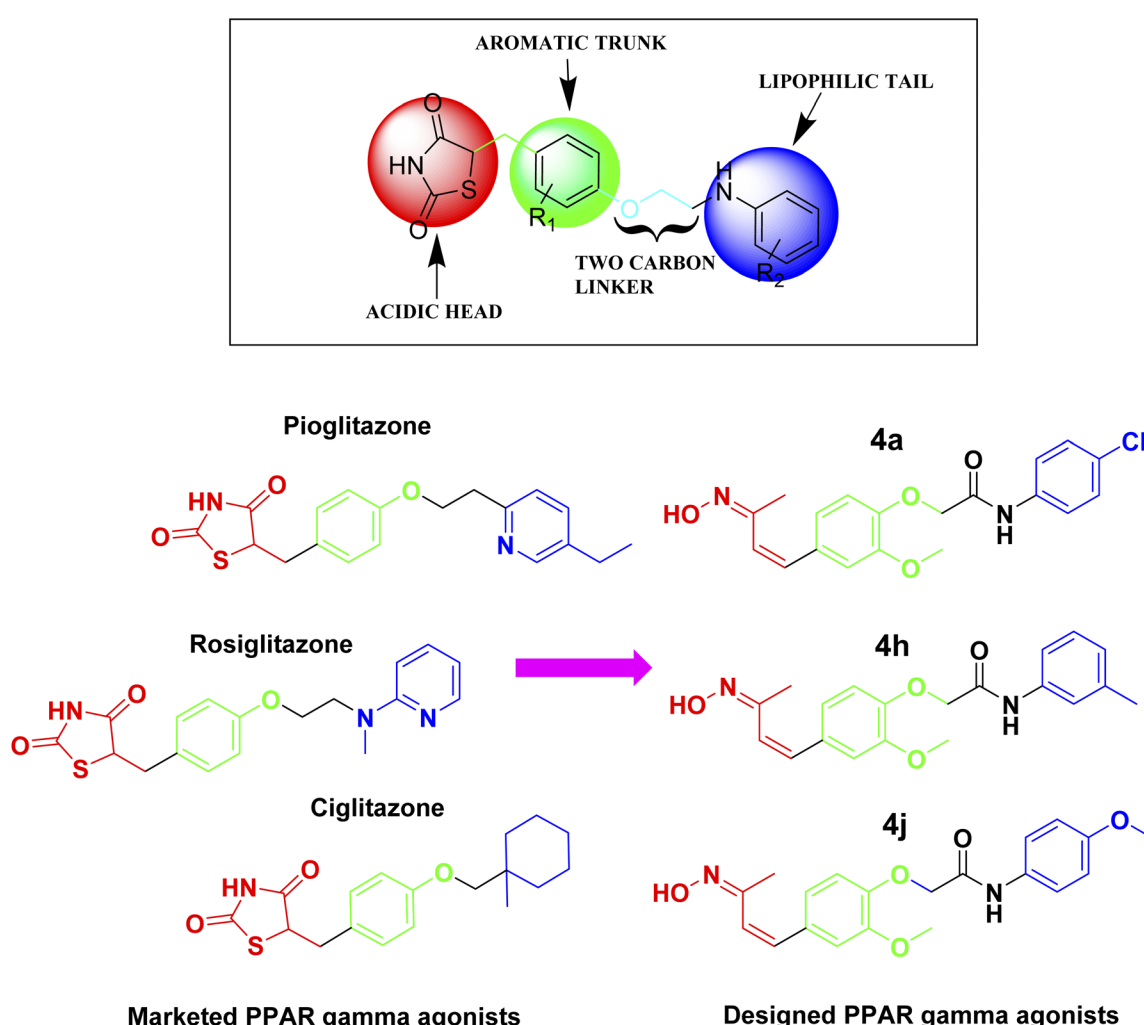


Fig. 1 Design strategy for the novel PPAR- γ agonists based on the structural features of available glitazones (PPAR- γ agonists).



Table 1 ADMET scores and toxicity profiles^a

Compd code	Solubility	BBB	HIA	CPY2D6	NTP_RAT		
					Male	Female	Ames mutagen
4a	3	2	0	NI	NC	NC	NM
4b	3	2	0	NI	NC	NC	NM
4c	3	3	0	NI	NC	NC	NM
4d	3	2	0	NI	NC	NC	NM
4e	3	2	0	NI	NC	NC	NM
4f	3	2	0	NI	NC	NC	NM
4g	4	3	0	NI	NC	NC	NM
4h	3	2	0	NI	NC	NC	NM
4i	3	2	0	NI	NC	NC	NM
4j	3	3	0	NI	NC	NC	NM

^a BBB: blood–brain barrier, HIA: human intestinal absorption, CYP2D6: cytochrome P450 2D6 (CYP2D6), NTP: national toxicology program.

health and cellular function. The development of synthetic ligands has allowed researchers to mimic the binding of ligands endogenously to activate PPAR- γ more effectively. Conditions such as AD, PD, ALS, MS, Huntington's disease, and stroke have all exhibited benefits from PPAR- γ ligand-induced receptor activation.²⁰

Therefore, after complete analysis of the structural features of available PPAR- γ agonists, novel PPAR- γ agonists were designed, synthesized, analyzed, and utilized in further *in vitro* studies. A TR-FRET binding assay was used to confirm the binding affinity of the synthesized compounds to PPAR- γ .²¹ Subsequent assessments were carried out to evaluate the *in vitro* cytotoxicity of these compounds. The results suggest that the development of novel PPAR- γ agonists with enhanced binding affinity and a predicted lower toxic profile holds promise for future therapeutic applications in AD. Notably, our studies demonstrated that novel PPAR- γ agonists have the potential to mitigate the production of pro-inflammatory cytokines, including TNF- α , IL-6, and IL-1 β , and also reduce oxidative stress in A β -induced SH-SY5Y neuroblastoma cell lines. This study delved into the potential binding interactions of these compounds with PPAR- γ using molecular docking and dynamics studies. Therefore, it becomes evident that the

activation of PPAR- γ receptors holds promise in regulating neuroinflammation and alleviating oxidative stress in the context of neurodegenerative diseases. In light of this, we undertook the design and synthesis of novel PPAR- γ agonists to explore their neuroprotective properties.

2. Results and discussion

2.1. Design rationale

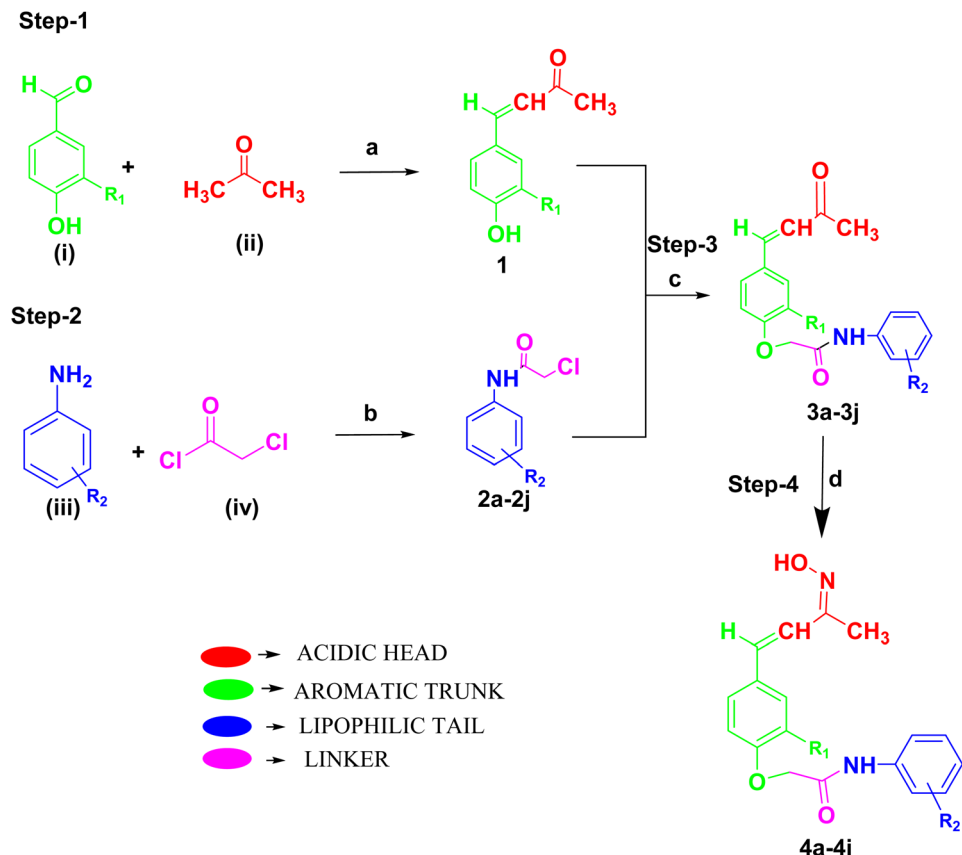
Glitazones are primarily used to manage type 2 diabetes due to their ability to improve insulin sensitivity, but some studies have also suggested glitazones may have potential benefits in AD.²² In the context of such research efforts, PPAR- γ agonists are known as glitazones that contain thiazolidinedione (TZD), a heterocyclic ring in the head region that is responsible for the activation of PPAR- γ receptor activation.²³ Our recent study aimed to design novel PPAR- γ agonists by replacing the TZD ring structure with a hydroxyl amine in the head region for PPAR- γ activation. With the understanding of the structural features of available glitazones, such as acidic head, aromatic trunk with two carbon linkers, and lipophilic tail, a newer focused library of molecules was designed (Fig. 1). Our study's approach of replacing the TZD ring with a hydroxyl amine in the

Table 2 Lipinski's rule of 5 and set dosage range for the rat model^a

Compound code	A log <i>p</i>	MW	HBA	HBD	Rat oral LD ₅₀ g per kg_body_weight	Rat inhalational LC ₅₀ mg m ⁻³ h ⁻¹	Carcinogenic potency TD ₅₀ _Rat mg per kg_body_weight per day
4a	3.363	374.82	6	2	2.55922	1361.53	17.3925
4b	3.185	354.41	6	2	0.92198	2000.58	544.362
4c	2.699	340.38	6	2	2.40968	1869.87	62.4133
4d	3.363	374.81	6	2	2.07843	1381.73	43.2746
4e	3.185	354.41	6	2	3.912	2107.68	114.23
4f	2.926	328.33	5	2	0.64346	2117.29	55.0068
4g	1.249	331.40	6	2	0.45388	867.721	150.922
4h	3.185	354.41	6	2	2.74511	2000.58	1.22823
4i	2.715	310.34	5	2	1.22902	1272.58	145.198
4j	2.682	370.39	7	2	4.63321	1289.51	1873.07

^a MW: molecular weight, HBA/D: hydrogen bond acceptor/donor.





Compound	R ₁	R ₂
4a	OCH ₃	<i>p</i> -Cl
4b	OCH ₃	<i>p</i> -CH ₃
4c	OCH ₃	-H
4d	OCH ₃	<i>o</i> -Cl
4e	OCH ₃	<i>o</i> -CH ₃

Compound	R ₁	R ₂
4f	H	<i>o</i> -F
4g	H	<i>p</i> -F
4h	OCH ₃	<i>m</i> -CH ₃
4i	H	-H
4j	OCH ₃	<i>p</i> -OCH ₃

Scheme 1 Synthesis: (a) NaOH, alcohol, and water with stirring for 6–7 h; (b) DCM/CHCl₃ and Et₃N, 0–5 °C to rt with stirring overnight; (c) K₂CO₃ anhydrous, KI, and acetone with stirring for 24 h; (d) sodium acetate and hydroxyl amine with stirring and reflux for 4 h.

head region for PPAR- γ activation is based on the concept of bioisosterism.²⁴ Bioisosteres are functional groups with a similar size, shape, and electron distribution, leading to comparable biological properties. In this case, the hydroxyl amine group might mimic the hydrogen-bonding interactions and polarity of the TZD ring, potentially maintaining PPAR- γ agonistic activity. A study by Saha *et al.*²⁵ supported this concept. They explored replacing the TZD ring in known drugs like rosiglitazone and pioglitazone with a pyrrolidinedione ring. Their findings showed that these analogs retained some PPAR- γ activity while exhibiting reduced toxicity compared to the parent TZD drugs. This highlights the potential for altering the heterocyclic ring structure while maintaining PPAR- γ agonism.

While specific examples of hydroxylamine-containing compounds with PPAR- γ activity are limited, the concept of using bioisosteric replacements is well-documented in medicinal chemistry. However, the replacement of TZD with a hydroxyl amine is a relatively unexplored bioisosteric approach. While there might not be documented examples yet, the rationale based on bioisosterism and the potential for hydrogen-bonding interactions justifies further investigation. Both functional groups can form hydrogen bonds, potentially interacting with similar binding sites on PPAR- γ . Our innovative structural features-guided design and synthesis of novel PPAR- γ agonists may offer a new perspective for this class of agonists for AD. The below graphical demonstration compares



the structural and functional attributes of the TZD ring (Original) and hydroxylamine moiety (Replacement), highlighting their respective roles in molecular interactions.

- TZD ring (Original):

- (1) Hydrogen bond donors and acceptors.
- (2) Electrostatic interactions.

(3) Planar structure facilitating binding.

- Hydroxylamine moiety (Replacement):

- (1) Hydrogen bond donors and acceptors.
- (2) Similar electronic properties.

(3) Potential to mimic planar interactions.

2.2. ADMET, TOPKAT, and drug likeness

Many drugs are out of the market due to their pharmacokinetics and dynamic properties, so this is a critical parameter that needs to be assessed at the beginning of the drug discovery cycle as a cost-effective and time-saving process.²⁶ By employing the small molecule tools in Discovery studio, the absorption-distribution-metabolism-excretion-toxicity (ADMET) scores and toxicity profiles²⁷ of compounds were calculated and are presented in Table 1. The compounds possessed good (level = 0) human intestinal absorption with good solubility (level = 3),^{28,29} except for compound 4g, which showed optimal solubility (level = 4). All the compounds were non-inhibitors of the metabolizing enzyme CYP450 activity^{30,31} and possessed a low (level = 3) to moderate (level = 2) ability to cross the blood-brain barrier (BBB). Furthermore, all the compounds obeyed Lipinski's rule of 5, which is a key descriptor for oral bioavailability, and potentially all the compounds were free from mutagenicity and carcinogenicity according to the rat oral LD50, inhalation LC50, and carcinogenic potency TD50 (ref. 32) indicators, as shown in Table 2.

Replacing the TZD ring with a hydroxylamine group in PPAR- γ agonists could improve their ADME properties, reduce toxicity, and enhance BBB penetration. Hydroxylamines can modulate lipophilicity for better absorption, improve metabolic stability, and decrease hepatotoxicity. They also have the potential to cross the BBB due to their favorable hydrophilic-lipophilic balance and molecular flexibility. Although specific examples of hydroxylamine-containing PPAR- γ agonists are limited, the principles of bioisosterism³³ and the existing literature on hydroxylamine derivatives support their potential. However, further empirical studies are needed to validate these theoretical benefits. There are reported studies that compounds containing hydroxylamine moieties are able to cross the BBB.³⁴ The designed ligands were studied for their favorable ADME properties, drug likeness, and toxicity, with a special focus on their ability to cross the BBB. The ability of a compound to penetrate the BBB can often be estimated by its log P value, which represents the compound's lipophilicity. A log P is useful for an initial assessment, and is one of Lipinski's rules to predict the logarithm of the partition coefficient (P) of a compound between octanol and water phases. It measures how well a compound can dissolve in lipids (octanol) *versus* water. To estimate the ability to cross the BBB, Egan and Lauri's method was employed based on the log P values, whereby:

1 Low log P (typically less than 2): compounds with very low log P values are usually too hydrophilic to efficiently cross the lipid-rich environment of the BBB.

2 Moderate log P (between 2 and 5): compounds with moderate log P values are often well-suited for BBB penetration. This range is considered optimal as it balances the hydrophilic and lipophilic properties necessary for crossing the BBB.

3 High log P (greater than 5): compounds with high log P values tend to be very lipophilic, which can result in poor aqueous solubility and increased non-specific binding to proteins and other components, potentially limiting their ability to penetrate the BBB effectively.

3. Chemistry and synthesis

Rationally designed compounds 2-(4-((1E,3E)-3-(hydroxylimino)but-1-en-1-yl)-2-methoxyphenoxy)-N-phenylacetamide derivatives (**4a–4j**) were synthesized according to Scheme 1. The structures of the synthesized compounds were confirmed using IR, ¹H NMR, ¹³C NMR, and mass spectral techniques. Four steps were employed to synthesize the final compounds. In the first step, two different aromatic aldehydes, such as 4-hydroxy-3-methoxy benzaldehyde (vanillin) and 4-hydroxy benzaldehyde, were (i) treated with acetone (ii) in the presence of NaOH to form the (E)-4-(4-hydroxy-3-methoxyphenyl)but-3-en-2-one derivative (**1**). In the second step, different aromatic amines (iii) were condensed with chloroacetyl chloride (iv) in the presence of triethylamine to form 2-chloro-N-(substituted phenyl) acetamide derivatives (**2a–2j**). The synthesized substituted phenols (**1**) and substituted phenyl acetamides (**2a–2j**) were reacted together in the presence of anhydrous K₂CO₃ and KI to form the prefinal compounds (*E*)-2-(4-(3-oxobut-1-en-1-yl)phenoxy)-N-(substituted phenyl) acetamide derivatives (**3a–3j**). The final compounds (**4a–4j**) were obtained by treating compounds **3a–3j** with hydroxylamine hydrochloride in the presence of sodium acetate. In Scheme 1, the final compounds (**4a–4j**) hydroxylamine (N=OH) were formed in the final step using hydroxylamine hydrochloride in the presence of sodium acetate and ethanol as a solvent under optimum temperature. The conditions provided in the synthesis of the titled compounds did not favor the formation of nitrosamine impurity. Therefore,

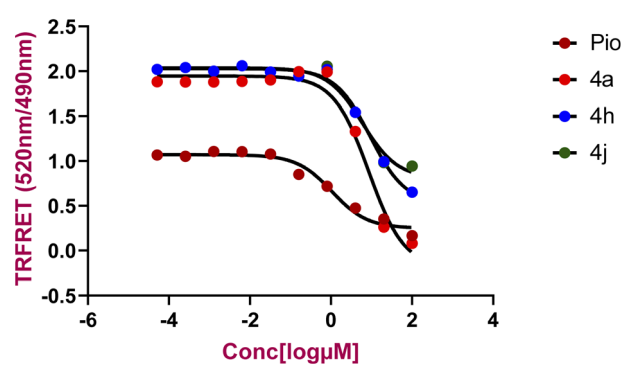


Fig. 2 Comparison of Lanthascreen TR-FRET PPAR- γ competitive binding assay results for pioglitazone, **4a**, **4h** and **4j** binding to the PPAR γ LBD.

we ruled out the possibility of the formation of nitroso amines as by-products in our synthesis of oximes. Moreover, the final synthesized compounds were purified by column chromatography and their formation was confirmed by checking their mass spectra. Then the structures were characterized and analyzed by NMR spectroscopy. The final compounds were confirmed by the appearance of ~ 1520.03 and $\sim 3313.11\text{ cm}^{-1}$ signals in the IR spectra, indicating C=N stretch and -OH stretch, respectively, while the appearance of a proton NMR peak at $\sim 11.01\delta$ ppm confirmed the -OH group formation, and an appropriate molecular ion peak (*M*) obtained by HRMS confirmed the formation of the final compounds.

4. PPAR gamma binding assay

For evaluation of the agonistic potential of the newly synthesized compounds, the Lanthascreen TR-FRET PPAR- γ competitive binding assay was employed. This assay pivots on the principle of competitive binding to PPAR- γ LBD, wherein the synthesized compounds displaced the fluorescent probe Fluromone. This displacement is quantifiable through a decrease in the 520/490 nm fluorescence ratio. The IC₅₀ values, representing the concentration required to displace 50% of Fluromone, were calculated from concentration-response curves, as presented in Fig. 2.

Among the tested compounds, **4a**, **4h**, and **4j** demonstrated binding to the PPAR- γ LBD, with IC₅₀ values of 8.60, 9.24, and 5.97 μM , respectively. For comparison, pioglitazone (Pio), a known PPAR- γ agonist, exhibited an IC₅₀ value of 1.052 μM represented in the Table 3. These findings indicate that the synthesized compounds exhibit a notable binding affinity to PPAR- γ LBD compared to pioglitazone.

5. In vitro studies

5.1. Cytotoxicity/cell viability assay

The synthetically derived compounds were assessed for their cytotoxicity against human neuroblastoma (SH-SY5Y) cells by MTT assay. The results of the cytotoxicity study performed by the MTT assay suggested that the compounds **4a** (108.4 μM), **4b** (111.8 μM), **4g** (121.5 μM), **4h** (150.0 μM), and **4j** (162.0 μM) were

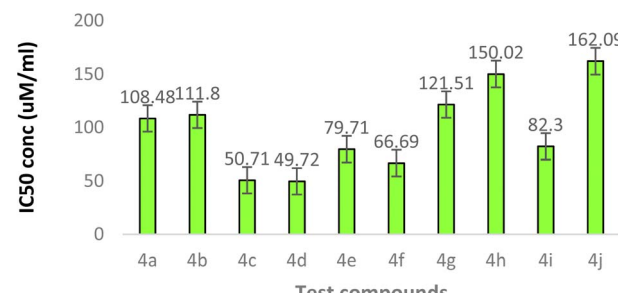


Fig. 3 MTT assay with IC₅₀ values of test compounds against SH-SY5Y cells.

cell proliferative and non-cytotoxic in nature with IC₅₀ values greater than 100 μM , while the other compounds **4c**, **4d**, **4e**, **4f**, and **4i** were cytotoxic in nature on SH-SY5Y cells with IC₅₀ values of 50.71, 49.72, 79.71, 66.69, and 82.30 μM , respectively (Fig. 3). The compounds that showed greater than 70% cell viability at the concentration of 50 μM were considered as cell proliferative against SH-SY5Y cells.

5.2. PPAR- γ transactivation assay

The PPAR- γ transactivation assay involved assessment of the particular transcription factor DNA binding activity in nuclear extracts. The findings showed that while all the compounds activated PPAR- γ in a cell-based transcription factor assay with a concentration range of 1–16 μM , notably, compounds **4a**, **4h**, and **4j** demonstrated significant biological efficacy, showing a dose-dependent enhancement in PPAR- γ transcriptional activity, as illustrated in Fig. 4. These compounds showed a significant expression of PPAR- γ in a concentration-dependent manner in SH-SY5Y cells that was comparable to that of pioglitazone, a typical full agonist. Based on these findings **4a**, **4h**, and **4j** are proposed as promising candidates for further inhibition assay to explore and optimize their therapeutic potential.

5.3. Anti-inflammatory and anti-oxidant activity in the A β -induced SH-SY5Y cell line

The anti-inflammatory and anti-oxidant activity of compounds **4a**, **4h**, and **4j**, against the A β -induced SH-SY5Y cell line were

Table 3 IC₅₀ values from PPAR- γ competitive binding assays of the synthesized compounds

Compd code	IC ₅₀ (μM)
4a	8.60
4b	17.25
4c	19.46
4d	13.27
4e	15.62
4f	14.03
4g	10.89
4h	9.24
4i	22.79
4j	5.97
Pio	1.05

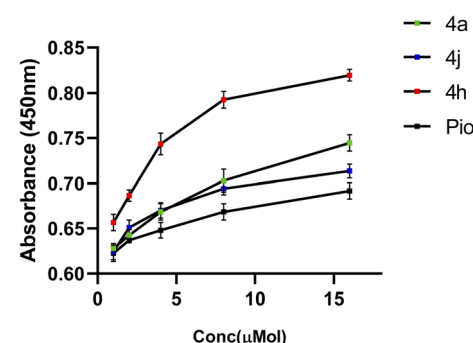


Fig. 4 PPAR- γ expression in SH-SY5Y cells treated with the test compounds (**4a**, **4h**, and **4j**) and pioglitazone at different concentrations (1–16 μM).



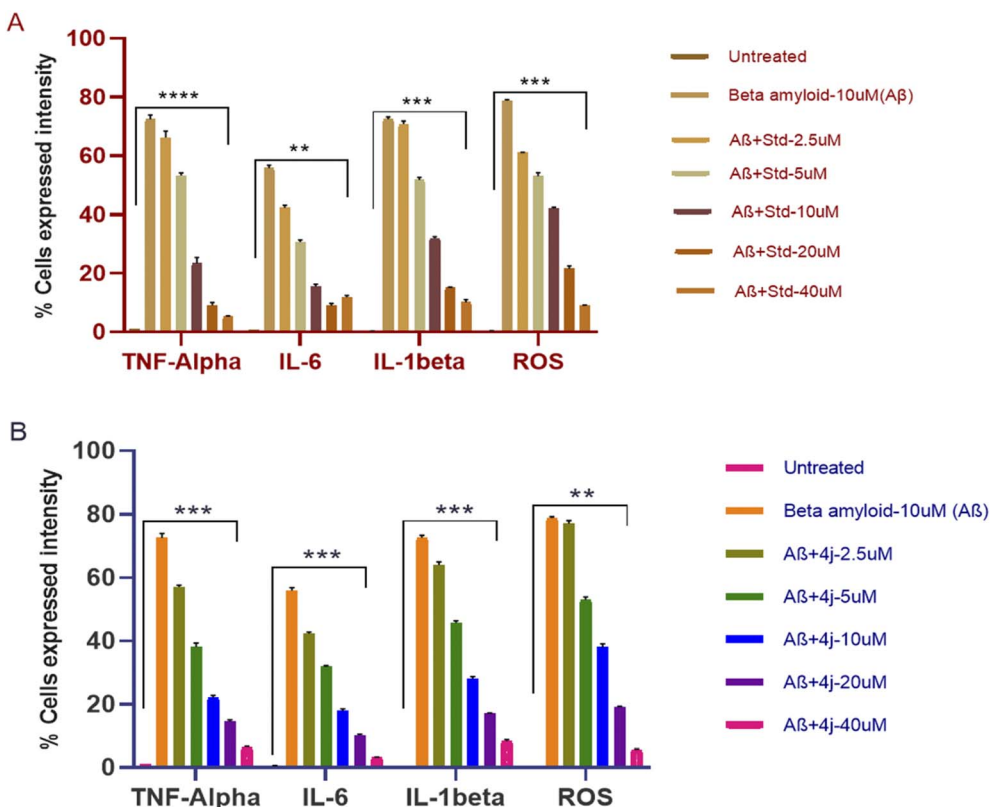


Fig. 5 (A and B) Bar graphs showing the % cell expression of TNF- α , IL-6 and IL-1 β and ROS in the A β -induced SH-SY5Y cell line upon treatment with pioglitazone and compound 4j. *Significance at p -value < 0.001 (***) and < 0.005 (**) compared to the untreated control.

analyzed by measuring the percentage expression of pro-inflammatory cytokines and ROS using flow cytometry. The compounds were taken in different concentration from 2.5 to 40 μ M. Most importantly, three of the synthesized compounds 4a, 4h, and 4j showed promising anti-inflammatory activity and anti-oxidant activity by reducing the expression of pro-inflammatory cytokines and ROS, respectively, in comparison to the standard (pioglitazone). The percentage decrease in the expression of cells by compound 4j showed its better activity in comparison to pioglitazone, as shown in Fig. 5.

5.3.1 Flow cytometric analysis of pro-inflammatory cytokines and ROS inhibition. In Table 4, “untreated” indicates that the SH-SY5Y cell line was not induced with A β and not treated

with the standard or the compounds. It was shown that there was a negligible percentage expression of cytokines and ROS in the normal SH-SY5Y cell line (1.05% TNF- α , 0.58% IL-6, 0.26% IL-1 β , and 0.37% ROS). The beta amyloid-10 μ M label indicates, the cell line was induced with only A β at a 10 μ M concentration to induce inflammation and stress mimicking AD conditions. The expressions for this factor were 73.52% TNF- α , 56.51% IL-6, 72.05% IL-1 β and 79.13% ROS. This indicates that there was an increase in the expression of pro-inflammatory cytokines and ROS, leading to neuroinflammation and neurodegeneration. Hence, the treatment was carried out with different concentration (2.5, 5, 10, 20, and 40 μ M) of the synthesized compounds 4a, 4h, and 4j, which were found to decrease the expression of

Table 4 The % cell expression of TNF- α , IL-6, IL-1 β cytokines and ROS in the A β -induced SH-SY5Y cell line by pioglitazone and compound 4j at different concentrations

Culture condition and concentration used	Pioglitazone				Compound 4j			
	TNF- α %	IL-6%	IL-1 β %	ROS%	TNF- α %	IL-6%	IL-1 β %	ROS%
Untreated	1.05	0.58	0.26	0.37	1.05	0.58	0.26	0.37
A β – 10 μ M (BAM)	73.52	56.51	72.05	79.13	73.52	56.51	72.05	79.13
B-AM + 2.5 μ M	67.8	42.95	70.21	60.98	41.44	48.78	48.22	77.72
B-AM + 5 μ M	53.92	30.23	52.38	53.96	35.32	36.58	45.00	53.58
B-AM + 10 μ M	24.85	15.09	32.36	41.78	19.75	26.69	26.02	38.78
B-AM + 20 μ M	9.8	9.51	15.15	21.01	15.35	23.22	16.4	18.82
B-AM + 40 μ M	5.5	12.31	10.76	9.07	8.43	6.52	4.88	5.53



pro-inflammatory cytokines and ROS; thereby exerting anti-inflammatory and anti-oxidant activities. Treatment with compound **4j** at the concentration of 10 μ M was found to decrease the expression of TNF- α by 19.75%, IL-6 by 26.69%, IL-1 β 26.02%, and ROS 38.78%. Similarly, pioglitazone at a 10 μ M concentration decreased the expression of TNF- α by 24.85%, IL-6 by 15.09%, IL-1 β by 32.36%, and ROS by 41.72%. These findings show that compound **4j** significantly decreased the expression of cytokines and oxidative stress when compared to the standard (pioglitazone). The decreases in the percentage expression of cytokines and ROS in a dose-dependent manner by the standard (pioglitazone) and compound **4j** are listed in Table 4.

The overlay plot of compound **4j** shows it reduced the markers associated with neuroinflammation and markers of stress in a dose-dependent manner compared to the untreated control cell line, as shown in Fig. 6. The histogram of compound **4j** is shown in Fig. 7, while the histogram of compounds **4a** and **4h** are given in the ESI (S:57–S:59).†

6. In silico evaluation

6.1. Molecular docking

The concept of structure-based drug design was implemented for known compounds and a drug target protein. The designed molecules were screened *in silico* by docking techniques using

modeling suite software, such as Discovery Studio (DS). The molecular docking was performed on PPAR- γ to understand the mode of ligand binding to the protein's active site and predict ligand potency to activate the protein in an agonistic sense. The synthesized molecules were docked against Pdb code: 3CS8 protein and the results were obtained in terms of -Cdocker energy and -C-Docker interaction energy. The interaction analysis between the receptor-ligand showed that all the compounds have the potential to bind at the ligand binding site, with varying docking energy scores listed in the Table 5. The -Cdocker energy was found to be significantly high in **4a**, **4h**, **4j** and in the standard (rosiglitazone). The ligand binding domain (LBD) regions of PPAR- γ amino acids were Lys261, Ile281, Cys285, Gln286, Ser289, Ile341, Met364, Lys367, His449, Leu453, Leu469, and Tyr473. Compounds **4a**, **4h**, and **4j** interact with active site amino acids Ile281, Cys285, Ser289, Ile341, Met364, Lys367, Leu453. A one or two hydrogen bond interaction was favored with the amino acids Ile326, Ile281, Ser 289, and Lys367 for compounds **4a**, **4h**, and **4j**, as well as pi–sulfur interaction with sulfur-containing amino acids at the active site, like Cys285 and Met364, while a hydrophobic interaction was observed with other active site amino acids. Common amino acids that interacted with the co-crystal ligand and the functional groups of the active molecules were Ile-341, Cys-285, Arg-288, Met-364, Leu-330, and Ala-292. These findings suggest that the compounds were successfully bound within the LBD of

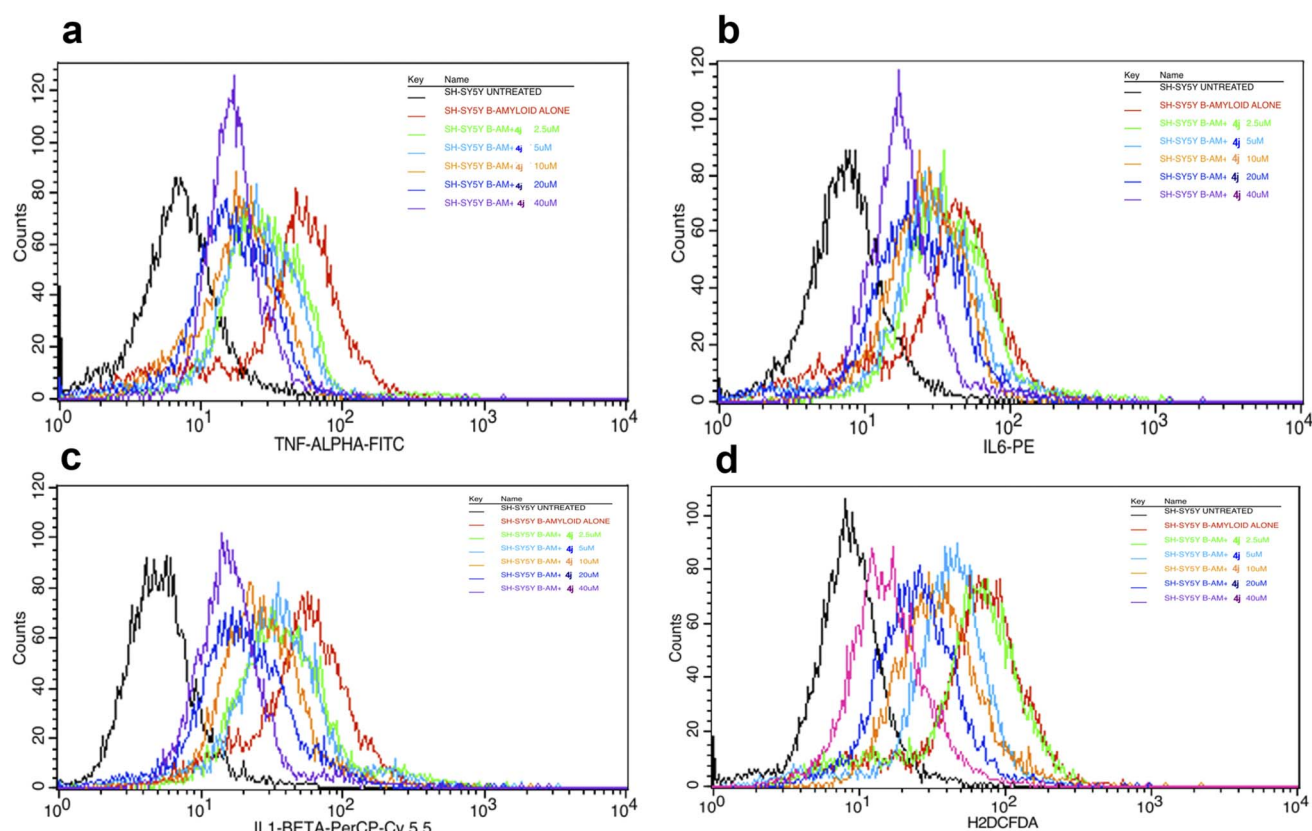


Fig. 6 Overlaid TNF- α (a), IL-6 (b), IL-1 β (c) and ROS (d) histograms in the A β -induced SH-SY5Y cell line for compound **4j** under untreated and treated conditions.



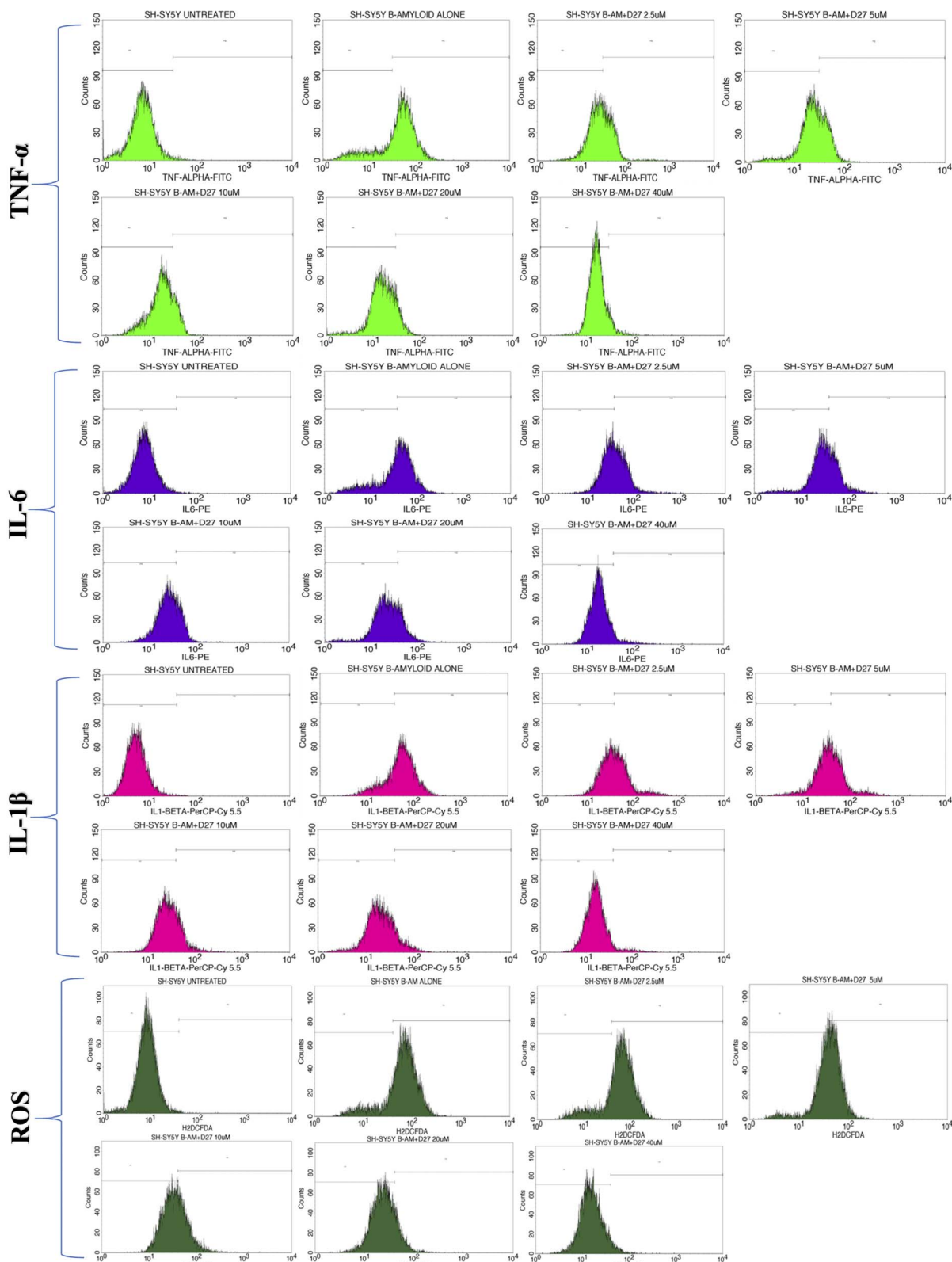


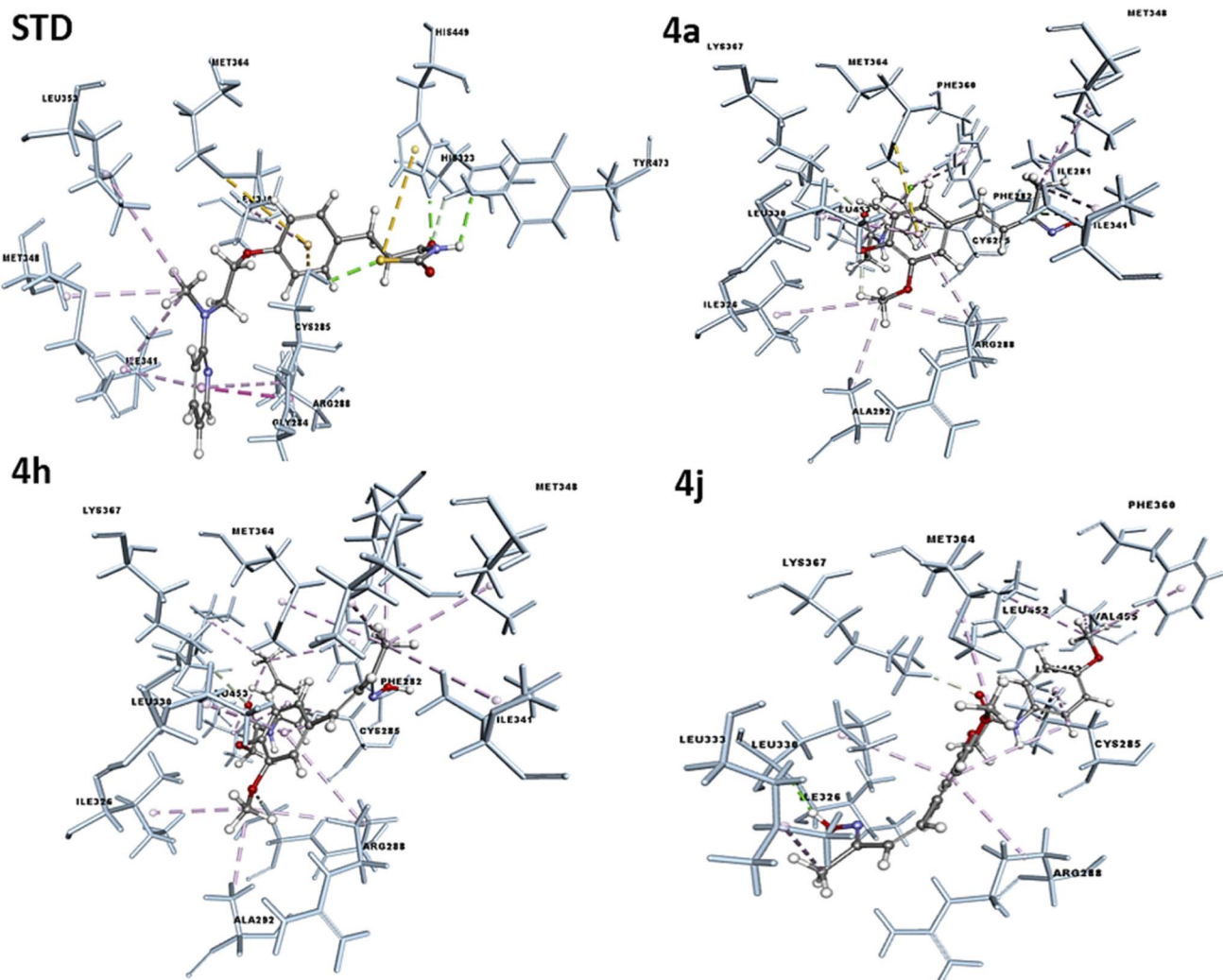
Fig. 7 Flow cytometry histogram showing the % cells expression of TNF- α , IL-6, and IL-1 β and ROS in the A β -induced SH-SY5Y cell line in different culture conditions, such as untreated, 10 μ M amyloid beta alone, and A β with compound 4j at different concentrations (2.5, 5, 10, 20, and 40 μ M) at the M1 and M2 phases. The intensity of peak in the M1 phase indicates there was a negligible expression of cells in the untreated condition. When the cell line was induced with A β , disease was induced and the % cells expression increased and the intensity of peak shifted to the M2 region. With the treatment of the induced cell line with compound 4j at concentrations of 2.5–40 μ M, the peak shifted towards the M1 region, which indicated the expression of cells decreased in a dose-dependent manner.

Table 5 Molecular dynamics-based docking with CDOCKER

Compounds	PPAR-gamma		
	-CDOCKER energy (kcal mol ⁻¹)	-CDOCKER interaction energy (kcal mol ⁻¹)	Potential interacting amino acids
4a	30.4483	46.7715	Ile281, Lys367
4b	27.4946	45.5273	Ser289, Met367
4c	21.4506	47.9125	Glu259, Met348
4d	27.2152	50.9619	Ser289, Leu330
4e	28.1738	41.5233	Leu453, Cys285
4f	28.5051	42.6889	His323, Ser342
4g	26.3025	44.6894	His323, Ile281
4h	29.3867	47.3176	Leu453, Ser289
4i	24.2704	45.7233	His323, Gly284
4j	31.6036	48.4746	Ile326, Lys367
Std (rosiglitazone)	36.3108	43.4691	Lys261, Met364

PPAR γ , with the optimum binding pattern shown in Fig. 8. All the investigated compounds demonstrated strong quantitative docking results compared to the standard (rosiglitazone) when docked with PPAR γ . The results of the docking study supported

the proposed hypothesis for the biological activity studies. The stability of these complexes was further analyzed by a molecular dynamic (MD) simulation study.

Fig. 8 Docking pose of the compounds **4a**, **4h**, **4j** and Std (rosiglitazone) to the active binding site of PPAR- γ .

6.2. Molecular dynamic studies

Molecular dynamics simulations of 50 ns duration were performed on 3CS8 protein complexed with compounds **4a**, **4h**, **4j** and the standard. These simulations aimed to characterize the receptor-ligand interactions over time. Subsequent analysis of the trajectories focused on the RMSD, RMSF, and protein-ligand interactions.³⁵

6.2.1 Root mean square deviation (RMSD). The RMSD plays a crucial role in analyzing the conformational changes in complex during dynamics. According to Dhivya *et al.*,³⁶ the level of confirmation is directly linked to the biological activity. Therefore, it is desirable to have lower deviations in RMSD. Overall, when comparing the synthesized compounds with the drug target PPAR- γ , the RMSD deviations remain within 4.5 Å. Initially, all the complexes showed a gradual increase in deviation, but after a few nanoseconds, complex **4j** demonstrated a deviation of 3.5 Å, and maintained stable equilibrium throughout the dynamics without exceeding 4.0 Å. Similarly, complex **4a** and the complex of the bound standard ligand exhibited similar trajectory paths with minimal deviations. However, complex **4h** exhibited a gradual increase in deviation, but remained stable within the deviation level of 4.5 Å. The RMSD plots are shown in Fig. 9.

6.2.2 Root-mean-square fluctuation (RMSF). RMSF measures the degree of fluctuation or movement of each residue in a biomolecular system. It helps identify the regions or residues that exhibit high flexibility or significant conformational changes. This information is crucial for understanding the structural dynamics of proteins and their functional implications. In drug discovery and design, studying the fluctuations in the active site of a protein is crucial. The active site residues of this drug target protein PPAR- γ included amino acids like Ile281, Cys285, Gln286, Ser289, Ile341, Met364, Lys367, His449, Leu453, Leu469, and Tyr473. The fluctuation of the active site

amino acids ranged between 0.5 Å and 1.2 Å, as shown in Table 6. Overall, the deviations of all the receptor-ligand complexes were within 3.0 Å, as shown in Fig. 10. Hence, by analyzing the RMSFs of the residues within the active sites, researchers can gain insights into their stability, conformational changes, and how they interact with potential ligands or inhibitors.

6.2.3 Radius of gyration (R_g). The R_g provides a measure of how compact or extended a biomolecule is in a given conformation. It quantifies the distribution of mass around the center of mass of the molecule. A smaller R_g value indicates a more compact and tightly folded structure, while a larger R_g value suggests a more extended or loosely packed conformation. Monitoring R_g during a simulation helps track changes in molecular compactness, which can provide insights into folding/unfolding events, structural transitions, or the effects of ligand binding. Overall, **4a**-3CS8, **4h**-3CS8, **4j**-3CS8, and Std-3CS8 complexes showed stable compactness throughout the dynamics and these complex deviations were between 18 Å and 19 Å without more deviations or changes in the trajectory path (Fig. 11), indicating the stability of the protein complexes and its relations to the complexes' compactness and structural integrity.

7. Structure–activity relationship

The structure–activity relationship (SAR)³⁷ is a fundamental consideration in organic chemistry that is instrumental in guiding the synthesis of compounds with enhanced potency for various drug targets. It helps predict the biological activity of compounds based on their chemical structure, with functional moieties playing a crucial role. In this study, receptor-ligand complexes (**4a**–**4j**) and their derivatives were generated using interaction pharmacophores for detailed analysis. A common 3-hydroxymino group in all compounds acted as a hydrogen

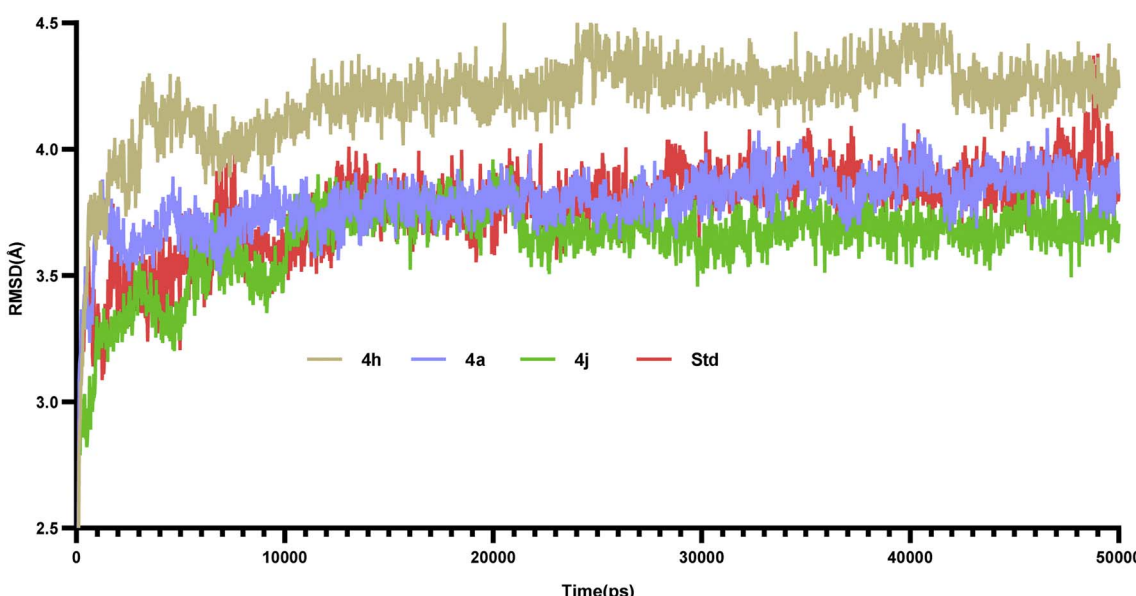
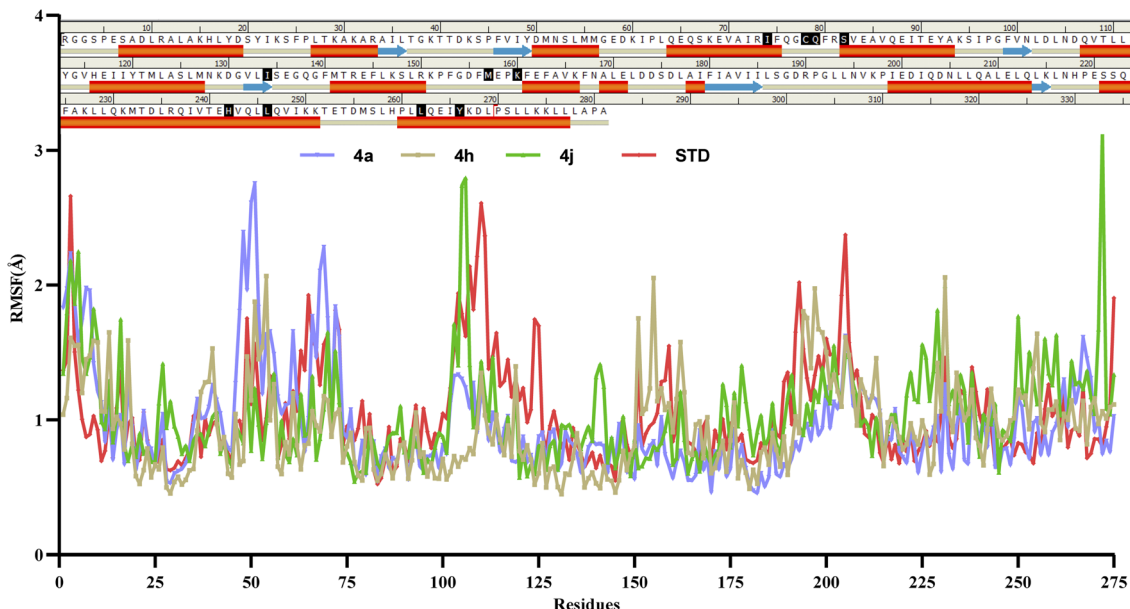


Fig. 9 RMSD plots of protein–ligand complexes of compounds **4a**, **4h**, **4j** and the Std (rosiglitazone).



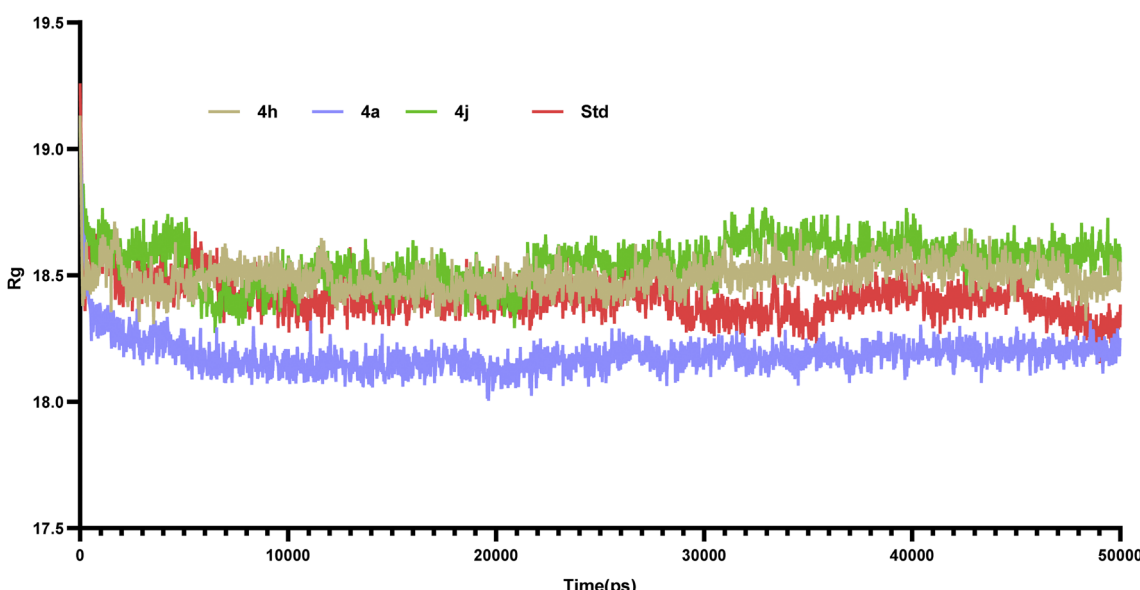
Table 6 Active site amino acids degree of fluctuations (Å)

Compounds	Active site amino acids (Å)										
	Ile281	Cys285	Gln286	Ser289	Ile341	Met364	Lys367	His449	Leu453	Leu469	Tyr473
4a	0.913	0.634	0.845	0.592	0.817	1.02	0.565	0.797	0.782	1.299	1.259
4h	0.687	0.589	0.552	0.937	0.718	1.233	0.91	0.926	0.849	1.045	0.921
4j	0.777	0.603	0.622	0.609	0.889	0.831	0.669	0.976	0.925	1.024	1.281
Std	0.814	0.934	1.14	0.801	0.941	1.283	0.841	1.227	0.785	1.209	0.885

Fig. 10 RMSF graph of protein–ligand complexes of compounds **4a**, **4h**, **4j** and the Std (rosiglitazone).

donor or acceptor, while the phenoxy trunk is vital for hydrophobic interactions. Notably, a compound at the lipophilic tail (benzene ring) with a different substitution pattern is critical for

mediating the hydrophobic interactions, which subsequently influences the activity. An optimal two-linker system is necessary to connect the phenoxy trunk and the lipophilic tail,

Fig. 11 R_g of protein–ligand complexes of compounds **4a**, **4h**, **4j** and the Std (rosiglitazone) over the entire time of simulation.

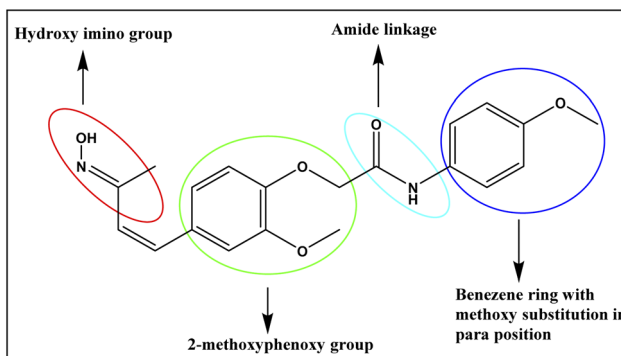


Fig. 12 Compound 4j's functional groups contributing to binding to the receptor.

ensuring these fragments fit into the active site pocket of PPAR- γ . The functional groups that contributes for the binding of compound to the active binding site of receptor was represented in the Fig. 12. The compound with a methoxy ($-\text{OCH}_3$) group at the *para* position (4j) demonstrated the highest activity among all the tested compounds. Similarly, the compound with a chloride (Cl) group at the *para* position (4a) also showed enhanced activity, whereas the same chloride substitution at the *ortho* position resulted in lower activity. Halogen substitutions, such as fluorine (F), at the *ortho/para* positions, showed slight deviations in activity, while a sole benzene ring exhibited minimal activity. A compound with a methyl (CH_3) group at the *meta* position (4h) showed optimal activity, whereas the *para* position substitution showed mild activity.

Hence, the functional groups in the benzene ring on the *para* and *meta* positions are crucial pharmacophores that can interact with the active site residues of drug targets, enhancing the anti-inflammatory effects in PPAR- γ binding and the expression of pro-inflammatory cytokines, as demonstrated by flow cytometry assays. Additionally, electron-donating groups, like methyl and methoxy, are also essential for the biological activity of these compounds.

8. Conclusion

In our study, based on the structural features of available PPAR- γ agonists, such as glitazones, we designed 10 novel compounds for the development of PPAR- γ agonists as neuroprotective agents through their anti-inflammatory and anti-oxidant properties. The synthesized compounds were characterized using analytical techniques. The new compounds were assessed for *in vitro* TR-FRET by the PPAR- γ competitive binding assay. Among all 10 synthesized compounds, 4a, 4h, and 4j showed significant binding affinity with the target protein compared with the standard drug. *In vitro* cytotoxicity evaluation of the all 10 compounds was carried with the SH-SY5Y human neuroblastoma cell line. Among all the other compounds, 4a, 4h, and 4j were cell proliferative and non-cytotoxic in nature, with IC_{50} values greater than 100 μM . The PPAR- γ agonistic activity of all the compounds was assessed using the Human PPAR- γ Transcription Factor Activity Assay Kit. Compounds 4a, 4h, and 4j were found to express PPAR-

γ in a concentration-dependent manner in the cell-based transcriptional factor assay. Compounds 4a, 4h, and 4j were assessed for their neuroprotective effect on the A β -induced SH-SY5Y cell line by measuring the intensity of reduction in proinflammatory cytokines and ROS by a flow cytometric method. The three compounds reduced the markers associated with neuroinflammation and markers of stress in a dose-dependent manner. Hence, we may infer that among the 10 new compounds, compounds 4a, 4h, and 4j showed the most promising activity by reducing neuroinflammation and reducing oxidative stress in the A β -induced cells, thereby exhibiting neuroprotection an effect by binding to PPAR- γ against AD. Furthermore, the molecular docking and MD simulation study results were correlated with the biological activity studies data, as the compounds with good binding interactions with PPAR- γ showed a promising effect on the SH-SY5Y cell line. Further analysis is required to clarify how these compounds ameliorate amyloid beta levels. Future pharmacodynamic and pharmacokinetic studies will be essential for advancing the application of these compounds. Hence, the findings of the current study suggest that the novel PPAR agonists exhibit potential neuroprotective properties, as inferred from their antioxidant and anti-inflammatory activities in the SH-SY5Y cell line under A β -induced stress. These findings support the potential use of novel PPAR- γ agonists in the treatment of Alzheimer's disease.

Repurposing marketed glitazones for AD treatment is a promising but complex endeavor. Glitazones activate PPAR- γ , influencing pathways relevant to AD pathogenesis, like glucose metabolism, inflammation, and oxidative stress.³⁸ Numerous preclinical studies have demonstrated the beneficial effects of glitazones in AD animal models.³⁹ These effects include improved cognitive function, reduced A β levels, and decreased neuroinflammation. Pioglitazone, in particular, has attracted attention due to its potential cognitive benefits in AD patients. Clinical trials have shown mixed results, with some indicating potential cognitive benefits but also highlighting safety concerns, such as cardiovascular risks and side effects, like weight gain.⁴⁰ The challenge lies in optimizing drug delivery to the brain, where effective concentrations are needed. Regulatory hurdles also exist, necessitating robust clinical evidence to support any new indication. While the evidence suggests that the drug repurposing of glitazones for AD treatment is a promising avenue, further research is needed to address the aforementioned challenges and fully realize the therapeutic potential of these compounds. A careful risk-benefit assessment is essential before their widespread clinical application in AD patients.

9. Experimental

9.1. ADMET, TOPKAT, and drug likeness

The pharmacokinetics and pharmacodynamics properties of the compounds were assessed through small molecular protocols (BIOVIA, Discovery Studio 2020) to understand the compounds behaviors. In addition, mutagenicity and carcinogenicity and the established dosage range of compounds were analyzed using a Bayesian model and regression model. Furthermore, the



Lipinski rule of 5 scores were calculated to determine the oral bioavailability of the compounds.⁴¹

9.2. Synthesis and chemistry

A novel series of PPAR- γ agonists were synthesized through the implementation of strategic synthetic methodologies. All the reactants, reagents, and solvents were procured from Sigma-Aldrich and utilized without further purification. The progress of the synthetic reactions was monitored *via* thin-layer chromatography (TLC), using precoated silica gel 60 F254 on aluminum plates. The synthesized compounds were purified through column chromatography.⁴² The reaction outcomes were further substantiated through spectroscopic analyses, including infrared (IR) spectroscopy, proton nuclear magnetic resonance ($^1\text{H-NMR}$), carbon-13 nuclear magnetic resonance ($^{13}\text{C-NMR}$), and mass spectrometry. The melting points of the synthesized compounds were determined using the open capillary method in a manual melting point apparatus, with the results reported in degrees celsius ($^{\circ}\text{C}$) without correction. Infrared spectra were recorded on a Shimadzu FT-IR 8400-S spectrometer employing the potassium bromide (KBr) pellet technique. Nuclear magnetic resonance (NMR) spectra (^1H and ^{13}C) were acquired using a Bruker 400 MHz FT-NMR spectrometer in deuterated dimethyl sulfoxide as a solvent, with the chemical shifts expressed in parts per million (ppm). Tetramethylsilane (TMS) was used as an internal standard. The coupling constants were recorded in Hertz (Hz). The HRMS of the synthesized molecules were determined utilizing ESI-MSMS (Make-Waters USS, Model-Xevo G2-XS Q TOF, Make-Waters, Illinois, IL, USA). For the analysis of the carbon, hydrogen, nitrogen, and sulfur (in the case of sulfur-containing compounds), an Elementar Vario (EL III Carl Erba11080) instrument was employed. The synthesis of the 10 novel PPAR- γ derivatives followed Scheme 1 and yielded compounds **4a–4j**. The TR-FRET binding assay kit (Catlog No. PV4894) was purchased from Invitrogen. Human neuroblastoma SHSY5Y cells were acquired from the National Centre for Cell Science (NCCS) in Pune, India. The pioglitazone was obtained from TCI Chemicals (CAS 11259-15-4) Lot QDIBK-NJ; purity > 98.0%.

9.2.1 General procedure for condensation of the substituted aromatic aldehyde with acetone (1). In a 250 mL flat-bottomed flask equipped with an ice bath, 0.1 M of acetone was mixed with 4 mL of 30% NaOH using a magnetic stirrer for 30 min. Then, to the mixture, 0.01 M of an aromatic aldehyde was added at room temperature. Next, 25 mL of ethanol was added to the reaction mixture, and stirred for 24 h. The reaction was checked for completion by TLC. Ice-cold water was added to the completed reaction mixture, which was then acidified with 10% HCl. The crude precipitate formed was filtered and washed with chilled water. The obtained crude product was subsequently purified by crystallization from a mixture of methanol and water.

9.2.2 General procedure for the synthesis of chloroacetylation with aromatic amine/heterocyclic amine compounds (2a–2j). The reaction mixture was prepared by the drop-wise addition of 2-chloroacetyl chloride (0.11 M) into

a solution containing different aryl amines/heterocyclic amines (0.1 M) and triethylamine (0.1 M) prepared in dichloromethane (80 mL) at 0 °C. The reaction mixture was brought to room temperature followed by stirring overnight. The reaction was checked for completion by TLC. Upon complete reaction, ice-cold water was added to the reaction mixture, which was then taken in a separating funnel and the organic layer was separated and then passed through anhydrous Na_2SO_4 . The organic layer was then distilled under pressure (Rota-evaporator) to obtain a crude product, which was subsequently purified by crystallization from a mixture of methanol and water.

9.2.3 General procedure for the condensation of acylated amines with substituted aromatic aldehydes (3a–3j). Acylated amine (0.01 M), substituted aromatic aldehyde (0.011 M), finely powdered anhydrous potassium carbonate (0.04 M), and potassium iodide (0.004 M) were dissolved in anhydrous acetone (30 mL). The reaction mixture was refluxed for 24 h and cooled to room temperature. The reaction was checked for completion by TLC. The solvent was removed under pressure and then ice-cold water was added to precipitate the crude product. The obtained crude product was subsequently purified by crystallization using ethanol.

9.2.4 General procedure for the synthesis of the final hydroxyimino series of compounds (4a–j). The reaction mixture was prepared by taking the synthesized pre final compounds from step-3 (3a–3j), *i.e.*, $\{(Z)\text{-N-(substituted phenyl)-2-(2-methoxy-4-(3-oxobut-1-en-1-yl)phenoxy)acetamide}\}$ (0.001 M), in a round-bottom flask containing 80 mL of ethanol. To this, hydroxylamine hydrochloride (0.006 M) in 10 mL of water and sodium acetate (0.01 M) in 10 mL of water were added. The reaction mixture was refluxed for 4 h in ethanol as a solvent. The reaction was checked for completion by TLC. The mixture was distilled under pressure to remove the solvent (ethanol). Ice-cold water was then added to reaction mixture with continuous stirring, and the crude product was precipitated out and was purified by column chromatography.

9.2.5 *N*-(4-Chlorophenyl)-2-(4-((1E,3E)-3-(hydroxyimino)but-1-en-1-yl)-2methoxyphenoxy)acetamide (4a). Light white amorphous solid, yield 60%, melting point 207–210, FTIR (KBr cm^{-1}): 3384 (N–H stretch), 3313 (O–H stretch), 2912 (C–H alkene stretch), 2851 (C–H alkane stretch), 1671 (C=O stretch), 1591 (N–H bending), 1520 (N–O stretch), 1238 (C–N stretch). $^1\text{H-NMR}$ (400 MHz, δ ppm, $\text{DMSO-}d_6$): 11.03 (s, 1H, $-\text{OH}$), 9.34 (s, 1H, NH), 8.07 (s, 1H, Ar–H), 7.59 (s, 1H, Ar–H), 7.27 (m, 3H, Ar–H, J = 8 Hz), 7.19 (d, 2H, Ar–H, J = 4 Hz), 7.17 (d, 1H, $\text{CH}=\text{CH}$, J = 20 Hz), 7.08 (d, 1H, $\text{CH}=\text{CH}$, J = 20 Hz), 4.87 (s, 2H, $-\text{OCH}_2$), 3.87 (s, 3H, $-\text{OCH}_3$), 2.20 (s, 3H, $-\text{CH}_3$). $^{13}\text{C-NMR}$ (100 MHz, δ ppm, $\text{DMSO-}d_6$): 27.84 (CH_3), 56.34 (OCH_3), 68.60 (OCH_2), 111.72 (1ArC), 114.24 (1ArC), 120.02 (2ArC), 123.23 (1ArC), 124.26 (1ArC), 126.27 (1ArC), 128.82 (1ArC), 129.40 (2ArC), 148.53 (1ArC), 138.98 (=CH), 143.85 (ArC) 149.86 (=CH), 150.16 (C=N), 166.83 (C=O). HRMS calculated [M + H] for $\text{C}_{19}\text{H}_{19}\text{ClN}_2\text{O}_4$, 375.1111; found [M + H] 375.1029. Anal. calc: C, 60.88; H, 5.11; Cl, 9.46; N, 7.47; found: C, 60.73; H, 5.12; Cl, 9.38; N, 7.41.

9.2.6 2-(4-((1E,3E)-3-(Hydroxyimino)but-1-en-1-yl)-2-methoxyphenoxy)-*N*-(*p*-tolyl)acetamide (4b). Dull white



amorphous solid, yield 70%, melting point 200–210, FTIR (KBr cm^{−1}): 3387 (N–H stretch), 3253 (O–H stretch), 3024 (C–H alkene stretch), 2935 (C–H aliphatic stretch), 1664 (C=O stretch), 1591 (N–H bending), 1551 (N–O stretch), 1249 (C–N stretch). ¹HNMR (400 MHz, δ ppm, DMSO-*d*₆): 11.04 (s, 1H, –OH), 9.95 (s, 1H, NH), 7.51 (d, 2H, Ar–H, J = 8 Hz), 7.23 (s, 1H, Ar–H), 7.14 (d, 2H, Ar–H, J = 8 Hz), 7.08 (d, 1H, Ar–H, J = 4 Hz), 6.95 (d, 1H, Ar–H, J = 8 Hz), 6.89 (d, 1H, CH=CH, J = 16 Hz), 6.80 (d, 1H, CH=CH, J = 16 Hz), 4.67 (s, 2H, –OCH₂), 3.86 (s, 3H, –OCH₃), 2.26 (s, 3H, –CH₃), 1.97 (s, 3H, –CH₃). ¹³CNMR (100 MHz, δ ppm, DMSO-*d*₆): 9.90 (CH₃), 20.90 (CH₃), 56.17 (OCH₃), 69.02 (OCH₂), 110.60 (1ArC), 114.92 (1ArC), 119.97 (2ArC), 120.28 (1ArC), 125.76 (1ArC), 129.61 (=CH), 131.24 (1ArC), 131.71 (2ArC), 133.11 (1ArC), 136.31 (=CH), 148.03 (1ArC), 149.85 (1ArC), 154.86 (C=N), 166.75 (C=O). HRMS calculated [M + H] for C₂₀H₂₂N₂O₄, 355.1658; found [M + H] 355.1596. Anal. calc: C, 67.78; H, 6.26; N, 7.90; found: C, 67.62; H, 6.31; N, 7.86.

9.2.7 2-(4-((1E,3E)-3-(Hydroxyimino)but-1-en-1-yl)-2-methoxyphenoxy)-*N*-phenylacetamide (4c). Off white amorphous solid, yield 65%, melting point 202–208, FTIR (KBr cm^{−1}): 3372 (N–H stretch), 3251 (O–H stretch), 3061 (C–H alkene stretch), 2935 (C–H alkane stretch), 1672 (C=O stretch), 1602 (N–H bending), 1553 (N–O stretch), 1251 (C–N stretch). ¹HNMR (400 MHz, δ ppm, DMSO-*d*₆): 11.04 (s, 1H, –OH), 10.04 (s, 1H, NH), 7.62 (s, 2H, Ar–H), 7.33 (m, 3H, Ar–H), 7.23 (s, 1H, Ar–H), 7.07 (s, 1H, Ar–H), 6.95 (d, 1H, CH=CH, J = 20 Hz), 6.85 (d, 1H, CH=CH, J = 20 Hz), 4.70 (s, 2H, –OCH₂), 3.86 (s, 3H, –OCH₃), 1.91 (s, 3H, –CH₃). ¹³CNMR (100 MHz, δ ppm, DMSO-*d*₆): 9.90 (CH₃), 56.17 (OCH₃), 68.98 (OCH₂), 110.60 (2ArC), 114.88 (1ArC), 119.25 (2ArC), 120.28 (1ArC), 124.14 (1ArC), 125.77 (1ArC), 129.25 (2ArC), 131.25 (1ArC), 131.71 (1ArC), 138.83 (ArC), 148.02 (=CH), 149.84 (=CH), 154.87 (C=N), 167.01 (C=O). HRMS calculated, [M + H] for C₁₉H₂₀N₂O₄ 341.1501; found [M + H] 341.1453. Anal. calc: C, 67.05; H, 5.92; N, 8.23; found: C, 67.15; H, 5.89; N, 8.26.

9.2.8 *N*-(2-Chlorophenyl)-2-(4-((1E,3E)-3-(hydroxyimino)but-1-en-1-yl)-2-methoxyphenoxy) acetamide(4d). White amorphous solid, yield 55%, melting point 208–210, FTIR (KBr cm^{−1}): 3387 (N–H stretch), 3308 (O–H stretch), 3064 (C–H alkene stretch), 2976 (C–H aliphatic stretch), 2918 (C–H stretch), 1681 (C=O) 1595 (N–H bending). ¹HNMR (400 MHz, δ ppm, DMSO-*d*₆): 10.99 (s, 1H, –OH), 9.29 (s, 1H, NH), 8.03 (s, 1H, Ar–H), 7.56 (d, 1H, Ar–H, J = 8 Hz), 7.21 (m, 3H, Ar–H, J = 8 Hz), 7.19 (d, 1H, Ar–H, J = 8 Hz), 7.14 (d, 1H, –CH=CH, J = 24 Hz), 7.06 (d, 1H, –CH=CH, J = 24 Hz), 6.98 (s, 1H, Ar–H), 4.72 (s, 2H, –OCH₂), 3.80 (s, 3H, –OCH₃), 2.18 (s, 3H, –CH₃). ¹³CNMR (100 MHz, δ ppm, DMSO-*d*₆): 17.86 (CH₃) 55.96 (OCH₃), 68.44 (OCH₂), 109.47 (2ArC), 114.32 (1ArC), 120.45 (2ArC), 124.27 (1ArC), 125.62 (1ArC), 126.59 (1ArC), 127.49 (1ArC), 130.80 (2ArC), 131.12 (1ArC), 135.97 (1ArC), 148.13 (=CH), 148.64 (=CH), 149.64 (C=N), 166.76 (C=O). HRMS calculated [M + H] for C₁₉H₁₉Cl N₂O₄, 375.1111; found [M + H] 375.1029. Anal. calc: C, 60.88; H, 5.11; N, 7.47; found: C, 60.46; H, 5.21; N, 7.78.

9.2.9 2-(4-((1E,3E)-3-(Hydroxyimino)but-1-en-1-yl)-2-methoxyphenoxy)-*N*-(*o*-tolyl)acetamide (4e). White amorphous solid, yield 75%, melting point 202–210, FTIR (KBr cm^{−1}): 3473

(N–H stretch), 3330 (O–H stretch), 2920 (C–H alkene stretch), 2856 (C–H alkane stretch), 1672 (C=O stretch), 1591 (N–H bending), 1549 (N–O stretch), 1264 (C–N stretch). ¹HNMR (400 MHz, δ ppm, DMSO-*d*₆): 10.43 (s, 1H, –OH), 9.35 (s, 1H, NH), 7.60 (d, 2H, Ar–H, J = 8 Hz), 7.41 (d, 1H, Ar–H, J = 4 Hz), 7.29 (m, 3H, Ar–H), 7.10 (t, 1H, Ar–H, J = 8), 7.085 (d, 1H, CH=CH, J = 16 Hz), 6.80 (d, CH=CH, J = 16 Hz), 4.67 (s, 2H, –OCH₂), 3.86 (s, 3H, –OCH₃), 2.26 (s, 3H, –CH₃), 1.97 (s, 3H, –CH₃). ¹³CNMR (100 MHz, δ ppm, DMSO-*d*₆): 9.90 (CH₃), 20.90 (CH₃), 56.17 (OCH₃), 69.02 (OCH₂), 110.60 (1ArC), 114.92 (1ArC), 119.97 (2ArC), 120.28 (1ArC), 125.76 (1ArC), 129.61 (=CH), 131.24 (1ArC), 131.71 (2ArC), 133.11 (1ArC), 136.31 (1ArC), 148.03 (=CH), 149.85 (=CH), 154.86 (C=N), 166.75 (C=O). HRMS calculated [M + H] for C₂₀H₂₂N₂O₄, 355.1658; found [M + H] 355.1558. Anal. calc: C, 67.78; H, 6.26; N, 7.90; found: C, 67.57; H, 6.32; N, 7.95.

9.2.10 *N*-(2-Fluorophenyl)-2-(4-((1E,3E)-3-(hydroxyimino)but-1-en-1-yl)phenoxy)acetamide (4f). Off white amorphous solid, yield 55%, melting point 208–212, FTIR (KBr cm^{−1}): 3385 (N–H stretch), 3271 (O–H stretch), 3037 (C–H alkene stretch), 2936 (C–H alkane stretch), 1676 (C=O stretch), 1596 (N–H bending), 1549 (N–O stretch), 1265 (C–N stretch). ¹HNMR (400 MHz, δ ppm, DMSO-*d*₆): 11.04 (s, 1H, –OH), 9.89 (s, 1H, NH), 7.84 (d, 1H, Ar–H, J = 4 Hz), 7.54 (d, 2H, Ar–H, J = 8 Hz), 7.31 (m, 4H, Ar–H), 7.00 (d, 2H, Ar–H, J = 8 Hz), 6.92 (d, 1H, CH=CH, J = 16 Hz), 6.76 (d, 1H, CH=CH, J = 20 Hz), 4.79 (s, 2H, –OCH₂), 1.982 (s, 3H, –CH₃). ¹³CNMR (100 MHz, δ ppm, DMSO-*d*₆): 9.91 (CH₃), 67.34 (OCH₂), 115.42 (2ArC), 115.98 (1ArC), 116.17 (1ArC), 124.87 (1ArC), 125.29 (1ArC), 125.71 (=CH), 125.83 (1ArC) 126.47 (1ArC), 126.83 (d, J _{C–F} = 7.1 Hz, C-3), 128.45 (2ArC), 130.32 (1ArC), 131.37 (=CH), 154.87 (C=N), 158.15 (C–F), 167.03 (C=O). HRMS calculated [M + H] for C₁₈H₁₇FN₂O₃, 329.1301; found [M + H] 329.1189. Anal. calc. C, 65.85; H, 5.22; N, 8.53; found: C, 65.72; H, 5.43; N, 8.65.

9.2.11 *N*-(4-Fluorophenyl)-2-(4-((1E,3E)-3-(hydroxyimino)but-1-en-1-yl)phenoxy)acetamide (4g). Greyish white amorphous solid, yield 55%, melting point 205–210, FTIR (KBr cm^{−1}): 3466 (N–H stretch), 3258 (O–H stretch), 3150 (C–H alkene stretch), 1664 (C=O stretch), 1579 (N–H bending), 1506 (N–O stretch), 1304 (C–N stretch). ¹HNMR (400 MHz, δ ppm, DMSO-*d*₆): 11.03 (s, 1H, –OH), 10.15 (s, 1H, NH), 7.68 (dd, 2H, Ar–H, J = 4 Hz), 7.54 (d, 2H, Ar–H, J = 12 Hz), 7.19 (t, 2H, Ar–H, J = 8 Hz), 7.01 (d, 2H, Ar–H, J = 8 Hz), 6.91 (d, 1H, CH=CH, J = 16 Hz), 6.75 (d, 1H, CH=CH, J = 16 Hz), 4.71 (s, 2H, –CH₂), 1.97 (s, 2H, –CH₂). ¹³CNMR (100 MHz, δ ppm, DMSO-*d*₆): 9.90 (CH₃), 68.70 (OCH₂), 115.46 (2ArC), 115.66 (2ArC), 122.02 (2ArC), 124.93 (d, J _{C–F} = 3.5 Hz, C-4), 125.36 (=CH), 128 (1ArC), 130.32 (1ArC), 131.237 (=CH) 135.18 (2ArC), 154.86 (C=N), 157.54 (1ArC–O), 159.93 (C–F), 166.86 (C=O). HRMS calculated [M + H] for C₁₈H₁₇FN₂O₃, 329.1301; found [M + H] 329.1189. Anal. calc: C, 65.85; H, 5.22; N, 8.53; found: C, 65.71; H, 5.36; N, 8.79.

9.2.12 2-(4-((1E,3E)-3-(Hydroxyimino)but-1-en-1-yl)-2-methoxyphenoxy)-*N*-(*m*-tolyl)acetamide (4h). Dull white amorphous solid, yield 80%, melting point 208–212, FTIR (KBr cm^{−1}): 3385 (N–H stretch), 3267 (O–H stretch), 2921 (C–H alkane stretch), 1666 (C=O stretch), 1615 (N–H bending), 1556 (N–O stretch), 1248 (C–N stretch). ¹HNMR (400 MHz, δ ppm, DMSO-*d*₆): 10.14 (s, 1H, –OH), 9.37 (s, 1H, NH), 7.48 (m, 3H, Ar–



H), 7.22 (d, 2H, Ar-H, J = 8 Hz), 7.03 (s, 1H, Ar-H), 6.94 (d, 1H, CH=CH, J = 16 Hz), 6.87 (d, 1H, CH=CH, J = 16 Hz), 6.41 (s, 1H, Ar-H), 4.73 (s, 2H, -OCH₂), 3.86 (s, 3H, -OCH₃), 2.09 (s, 3H, -CH₃), 2.01 (s, 3H, -CH₃). ¹³CNMR (100 MHz, δ ppm, DMSO-*d*₆): 9.90 (CH₃), 21.63 (CH₃), 56.12 (OCH₃), 68.84 (OCH₂), 110.47 (1ArC), 114.67 (1ArC), 117.08 (1ArC), 120.41 (2ArC), 124.82 (=CH), 125.70 (1ArC), 129.10 (1ArC), 131.15 (=CH), 131.74 (1ArC), 138.48 (1ArC), 138.76 (1ArC), 147.99 (1ArC), 149.76 (1ArC), 154.89 (C=N), 166.93 (C=O). HRMS calculated [M + H] for C₂₀H₂₂N₂O₄, 355.1658; found [M + H] 355.1596. Anal. calc. C, 67.78; H, 6.26; N, 7.90; found: C, 67.82; H, 6.17; N, 7.89.

9.2.13 2-(4-((1E,3E)-3-(Hydroxyimino)but-1-en-1-yl)phenoxy)-N-phenylacetamide (4i). Yellowish white amorphous solid, yield 60%, melting point 200–208, FTIR (KBr cm⁻¹): 3392 (N-H stretch), 3196 (O-H stretch), 3060 (C-H alkene stretch), 2918 (C-H alkane stretch), 1660 (C=O stretch), 1618 (N-H bending), 1538 (N-O stretch), 1247 (C-N stretch). ¹HNMR (400 MHz, δ ppm, DMSO-*d*₆): 10.15 (s, 1H, -OH), 9.31 (s, 1H, NH), 7.68 (dd, 2H, Ar-H, J = 8 Hz), 7.50 (d, 2H, Ar-H, J = 8 Hz), 7.19 (t, 1H, Ar-H, J = 8 Hz), 7.02 (d, 2H, Ar-H, J = 8 Hz), 6.92 (d, 1H, CH=CH, J = 16 Hz), 6.78 (d, 1H, CH=CH, J = 16 Hz), 6.38 (s, 1H, ArH), 4.72 (s, 2H, -CH₂), 2.00 (s, 2H, -CH₂). ¹³CNMR (100 MHz, δ ppm, DMSO-*d*₆): 11.98 (CH₃), 67.61 (OCH₂), 115.53 (2ArC), 122.01 (2ArC), 128.26 (2ArC), 130.39 (=CH), 131.12 (1ArC) 135.18 (=CH), 146.12 (1C, ArC-N), 157.42 (C=N), 158.20 (1C, ArC-O), 166.86 (C=O) 1.97 (s, 3H, -CH₃). HRMS calculated [M + H] for C₁₈H₁₈N₂O₃, 311.1395; found [M + H] 311.1589. Anal. calc. C, 69.66; H, 5.85; N, 9.03; found: C, 69.71; H, 5.84; N, 9.11.

9.2.14 2-(4-((1E,3E)-3-(Hydroxyimino)but-1-en-1-yl)-2-methoxyphenoxy)-N-(4-methoxyphenyl) acetamide (4j). White amorphous solid, yield 80%, melting point 202–208, FTIR (KBr cm⁻¹): 3384(N-H stretch), 3196 (O-H stretch), 3016 (C-H alkene stretch), 2838 (C-H alkane stretch), 1653 (C=O stretch), 1608 (N-H bending), 1556 (N-O stretch), 1247 (C-N stretch). ¹HNMR (400 MHz, δ ppm, DMSO-*d*₆): 11.04 (s, 1H, -OH), 9.90 (s, 1H, NH), 7.54 (d, 2H, Ar-H, J = 9.2 Hz), 7.23 (d, 1H, Ar-H, J = 1.6 Hz), 7.06 (d, 1H, CH=CH, J = 2 Hz), 6.95 (m, 3H, Ar-H), 6.89 (d, 1H, CH=CH, J = 16 Hz), 6.80 (d, 1H, CH=CH, J = 16 Hz), 4.66 (s, 2H, -OCH₂), 3.86 (s, 3H, -OCH₃), 3.73 (s, 3H, -OCH₃), 1.98 (s, 3H, -CH₃). ¹³CNMR (100 MHz, δ ppm, DMSO-*d*₆): 9.59 (CH₃), 55.98 (OCH₃), 56.12 (OCH₃), 66.92 (OCH₂), 111.52 (1ArC), 115.34 (2ArC), 122.56 (3ArC), 124.36 (=CH), 127.26 (1ArC), 130.81 (1ArC), 137.32 (=CH), 147.35 (1ArC), 149.75 (1ArC), 156.24 (1ArC), 156.28 (C=N), 158.91 (1ArH), 167.91 (C=O). HRMS calculated [M + H] for C₂₀H₂₂N₂O₅, 371.1607; found [M + H] 371.1495. Anal. calc: C, 64.85; H, 5.99; N, 7.56; found: C, 64.76; H, 5.95; N, 7.68.

9.3. TR-FRET assay

The study investigated the ability of the synthesized compounds to interact with the PPAR- γ ligand binding domain (PPAR- γ -LBD) using the Lanthascreen TR-FRET PPAR γ competitive binding assay kit from Invitrogen. Following the instructions provided with the kit, the procedure involved mixing 0.5 nM of GST-tagged human PPAR- γ -LBD, 5 nM of terbium-labeled anti-GST antibody, 5 nM of a fluorescent PPAR ligand (Fluormone

Pan-PPAR Green), and varying concentrations of the new compounds along with pioglitazone. This mixture was then incubated in the dark for 2 to 3 h. The FRET signal was detected using a Varioskan LUX multimode microplate reader (Thermo Fischer) by exciting at 340 nm and recording emissions at 530 nm for FluormoneTM Pan-PPAR Green and 490 nm for terbium. The interaction with PPAR- γ -LBD was assessed based on the ratio of the emission at 530 nm and 490 nm.⁴³ The principle behind this is that the test compounds, when binding to the PPAR- γ -LBD, will displace the fluorescently labeled ligand, leading to a reduced FRET signal. This decrease in the FRET signal, indicated by the 530 nm/490 nm ratio, can serve as an indicator of the compound's binding efficiency to the human PPAR γ LBD.⁴⁴ The results were processed using Prism software (GraphPad Software 08) and the sigmoidal curve equation with a variable slope was applied to determine the IC₅₀ values of the compounds.

9.4. Cytotoxicity/cell viability assay

For the MTT assay conducted on the neuroblastoma (SH-SY5Y) cell line, cells were cultured under standard conditions and seeded in a 96-well plate at a density optimized for growth, typically ranging from 5000–10 000 cells per well. Following 24 h adherence, the cells were treated with various concentrations of the synthesized compounds and the controls comprising untreated cells. Post 24–72 h of incubation, the cell viability was assessed using the MTT assay.^{45,46} This involved replacing the treatment media with a fresh MTT solution (5 mg mL⁻¹ in PBS), incubating for 2–4 hours to allow for formazan crystal formation, followed by solubilization using DMSO or isopropanol. The absorbance was measured at 570 nm using a microplate reader, correlating directly with living cell numbers. Data were analyzed by comparing treated and untreated cells' absorbance values to calculate the IC₅₀ value. The IC₅₀ value was determined by using a linear regression equation. The percentage cell viability was calculated using the below formula.

$$\% \text{ cell viability} = \left[\frac{\text{mean abs of treated cells}}{\text{mean abs of untreated cells}} \right] \times 10$$

9.5. PPAR-gamma transactivation assay

The activity of PPAR- γ transcription factor was evaluated with SH-SY5Y cell line nuclear extract using a sensitive ELISA-based method.⁴⁷ This assay could specifically measure the DNA-binding activity of ligand-activated PPAR- γ in nuclear extracts from these cells. Following the manufacturer's protocol for the Human PPAR- γ Transcription Factor Activity Assay Kit [TFEH-PPAR γ -1 (Ray Biotech, GA)], we determined the activity levels of these transcription factors by quantifying the absorbance at 450 nm. In the experimental setup, SH-SY5Y cells were first exposed to the compounds under investigation. Subsequently, nuclear proteins were isolated from the treated cells using Ray Biotech's Nuclear Extraction Kit (Cat No: NE50). The extraction process involved pelleting the nuclear contents through centrifugation at 15 000 rpm for 10 min at a temperature of 4 °C.



This methodology allowed for the precise assessment of PPAR- γ transcription factor activities in response to the test compounds.⁴⁸

9.6. Pro-inflammatory and ROS profiling in A β -induced SH-SY5Y cells

9.6.1 Flow cytometric analysis for TNF alpha, IL-6 and IL-1 beta expression inhibition. In this study, we measured the levels of the pro-inflammatory cytokines TNF- α , IL-6, and IL-1 β in a SH-SY5Y cell line induced with A β using flow cytometry. The cells were first cultured and then exposed to A β to induce inflammation. After allowing sufficient time for cytokine production, we fixed and permeabilized the cells to enable antibody staining. We then used specific antibodies tagged with the fluorescent markers TNF- α FITC, IL-6 PE, and IL-1 β PerCP Cy5.5 to identify the expression of TNF- α , IL-6, and IL-1 β respectively. The flow cytometry technique was utilized to analyze the fluorescence, which indicated the amount of each cytokine present in the cells. Our analysis focused on comparing the treated cells with untreated controls to determine the cytokine expression levels.

9.6.2 Flow cytometric analysis of ROS expression inhibition. Next, we evaluated the levels of reactive oxygen species (ROS) in the SH-SY5Y cell line after A β induction, employing flow cytometry. Primarily, the selected neuronic cell line was cultured under standard conditions and then treated with A β to induce oxidative stress. To measure ROS, cells were incubated with a ROS-sensitive fluorescent dye, such as 2',7'-dichlorofluorescin diacetate (DCFDA), which produces fluorescence upon reacting with ROS. After the incubation period, the cells were washed and analyzed using flow cytometry. This method enabled us to detect and quantify the fluorescence, which was directly consistent with the ROS levels in the cells. Control samples were included to provide a reference point for assessment.

9.7. Molecular docking

For the molecular docking, the structures of the drug target receptor (PDB ID:3CS8)⁴⁹ and compounds were prepared to mimic the *in vivo* environment. Both receptor and ligands were prepared using the Prepare Protein and Prepare Ligands protocol in Discovery Studio 2020. Further, the receptor binding site was defined on the current selection of the bound ligand coordinates 25.025X 1.3175Y 27.177Z for a radius of 7.9 Å and equal grid spacing of 0.5 Å with 90-degree grid angles. Molecular dynamic grid-based docking CDOCKER was used for docking the multiple compounds at the defined site of binding.⁵⁰ Before docking, validation of re-docking of the native bound compound was required for parameter optimization to obtain the docked poses within 2.5 Å. Finally, all the compounds were docked to the modified CDOCKER protocol. The best pose of ligands was taken for the molecular dynamics simulation and for the non-bonded interaction analysis.

9.8. Dynamics and simulation

To assess the stability and real-time behavior of the PPAR- γ docked complex dynamics, a CHARMM simulation was conducted for 50 000 ps. The BIOVIA Discovery Studio suite was utilized, implementing the CHARMM force field parameters for four complexes, namely **4a**-3CS8, **4h**-3CS8, **4j**-3CS8, and Std-3CS8. The protein-ligand complex systems were solvated using the TIP3 water model in an orthorhombic box and neutralized with 0.145 M NaCl. Two stages of energy minimization were performed, consisting of 500 steps each using the steepest descent and conjugate gradient algorithms. Subsequently, the system was gradually heated, equilibrated, and subjected to production under the NPT isothermal-isobaric ensemble. Temperature control was achieved using the Nose-Hoover thermostat, maintaining a temperature of 300 K, while the pressure control was set at 1.0 bar using the Langevin piston method. The dynamics Integrator employed was the Leapfrog Verlet Integrator, enabling numerical integration of the equation of motion. To assess the stability of the complex, time-dependent parameters, such as RMSD (root-mean-square deviation), RMSF, and the radius of gyration, were calculated.^{51,52}

Abbreviations

TLC:	Thin-layer chromatography
IC ₅₀ :	Inhibitory concentration 50
BBB:	Blood-brain barrier
NI:	Non-inhibitor
NC:	Non-carcinogen
NM:	Non-mutagen
DMSO:	Dimethyl sulfoxide
RO5:	Lipinski's rule of 5
HBD:	Hydrogen bond donor
HBA:	Hydrogen bond acceptor
HIA:	Human intestinal absorption
TR-FRET:	Time-resolved fluorescence energy transfer
ELISA:	Enzyme-linked immunosorbent assay
TNF- α :	Tumor necrosis factor-alpha
IL-6:	Interleukin 6
IL-1 β :	Interleukin 1 beta
ROS:	Reactive oxygen species
RXR:	Retinoid X receptors

Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

Conceptualization, B. R. P. K. and P. D.; methodology, P. D.; software, D. S and P. D.; analysis and validation, B. R. P. K. and P. P.; resources and data curation, P. D.; writing original manuscript, P. D.; writing – review and editing, P. D., A. P.; visualization and formal analysis, B. R. P. K., U. H., N. H., S. T.;



guidance, B. R. P. K.; funding acquisition, U. H., N. H., S. T.; project administration, B. R. P. K.

Conflicts of interest

The authors declare no financial conflicts of interest or personal affiliations that could have influenced the research conducted in this study or any kind of conflict of interest.

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