MINI REVIEW ON THERAPEUTIC PROFILE OF PHENOXY ACIDS AND THEIR DERIVATIVES

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ABSTRACT
Phenoxy acids and their derivatives are associated with a variety of biological activities such as antihyperlipidemic, hypoglycemic, antimicrobial, antiviral, antitubercular, anti-inflammatory, analgesic, antioxidant, anticancer and antihypertensive activities. This mini review outlines diverse biological properties of phenoxy acids and their derivatives.

Keywords: Phenoxy acids, Antitubercular, Anti-inflammatory, Analgesic, Antioxidant

INTRODUCTION
The phenoxy acetic acid moiety is present in diverse pharmacological agents such as hypolipidemic, analgesic, anti-inflammatory, antitubercular, antibacterial and antifungal agents.

Hypolipidemic activity
Fenofibrate (1) and gemfibrozil (2), clinically available hypolipidemic agents possess phenoxy propionic acid moiety in their structure and found to act as weak agonists of peroxisome proliferator-activated receptor alpha (PPAR-α). Hyperlipidemia, characterized by increased blood cholesterol and triacylglycerol levels is considered as the principal risk factor for several life-threatening cardiovascular diseases [1]. These drugs are highly effective in treating hyperlipidemia by increasing high-density lipoproteins (HDL) levels.

Phenoxy acids and their derivatives containing various heterocyclic rings exhibited significant hypolipidemic activity in different animal models [2].

Li et al. synthesized novel phenoxy alkyl carboxylic acid derivatives (fig. 3) based on the natural scaffolds (flavonoids or resveratrol) and evaluated them for hypolipidemic activity using triton WR-1339-induced hyperlipidemia and alloxan-induced diabetic models. Derivative bearing resveratrol scaffold lowered the triglyceride levels by 48.5%, total cholesterol by 44.2% and was active comparable to that of the standard drug fenofibric acid in the triton-induced hyperlipidemic model when administered orally at a dose of 300 mg/kg body weight. This derivative significantly lowered triglycerides levels in an alloxan-induced diabetic model upon oral administration at a dose of 150 mg/kg body weight. Molecular
docking studies revealed that the most active compound can bind well with the target enzyme PPAR-α [4].

![Fig. 3](image)

### Antinociceptive and Antiinflammatory activity

A series of novel phenoxy acid hydrazide derivatives containing carboxylic moiety such as 4-oxobutanoic acid at 4th position (fig. 4) were synthesized, and their antinociceptive and anti-inflammatory activities were evaluated. Most of the derivatives were found to exhibit peripheral nociceptive effects. The results demonstrated that the presence of acidic moiety increased the peripheral antinociceptive activity and reduced the central antinociceptive activity [5].

![Fig. 4](image)

Gheraldini et al. synthesized phenoxy ester derivatives containing N-methylpropane ring such as SM 21, (3-α-tropanyl) 2-[4-Cl-phenox] butyrate and PG-9, α-tropanyl 2-[4-bromophenyl] propionate (fig. 5) keeping in view that atropine induces central antinociception by increasing cholinergic transmission. These derivatives were evaluated for their central and peripheral analgesic activities in different animal models and nootropic properties. Both the compounds showed promising analgesic activities and also exhibited nootropic activities [6, 7].

![Fig. 5](image)

A novel series of 2-(substituted phenoxy)-N-(1-phenylethyl) acetamide derivatives (fig. 6) were synthesized using 1-phenylethylamine and substituted phenols. The synthesized amides were evaluated for their anticancer activity against MCF-7 (breast cancer), SK-N-SH (neuroblastoma), anti-inflammatory activity and analgesic activity. The results showed that halogen-containing derivatives increased the anticancer and anti-inflammatory activity whereas nitro group bearing derivatives exhibited good anticancer, anti-inflammatory and analgesic activities [8].

![Fig. 6](image)

Bhavna et al. synthesized phenoxy ester and amide derivatives (fig. 7) using paracetamol. The synthesized compounds showed promising analgesic and antipyretic activities comparable to the standard drug paracetamol [9].

![Fig. 7](image)

### Hypoglycemic activity

Nikalje et al. synthesized a series of novel phenoxy amide derivatives possessing 2, 4-thiazolidinedione moiety (fig. 8) by mixing 2, 4-thiazolidinedione and 2-(4-formyl phenoxy) N-substituted acetamide at room temperature. The synthesized compounds were evaluated for their hypoglycemic activity in mice model and the results showed that several derivatives bearing electron-releasing substituents such as methyl or methoxy groups exhibited promising hypoglycemic activity [10].

![Fig. 8](image)

The same research group further synthesized a series of derivatives using 2-methoxyphenoxy-N-substituted acetamide (fig. 9) and reported significant hypoglycemic activity with all the derivatives. Good activity was observed with derivatives containing electron releasing methoxy group on phenyl ring bearing phenoxy acid scaffold [11].

![Fig. 9](image)

### Anticancer and antiviral activity

Shahar Yar et al. synthesized substituted phenoxy acetic acid derived pyrazolines (fig.10) by the reaction between 2-{4-[3-(2, 4-dihydroxyphenyl)-3-oxo-1-propenyl]-2-methoxyphenoxy} acetic acid and various acid hydrazides and screened them for their in vitro cytotoxicity and antiviral activity.

![Fig. 10](image)
The highest cytotoxic activity was observed with 2-hydroxy phenyl derivative against human embryonic lung (HEL) cells, at a minimum cytotoxic concentration (MIC) of 0.16 µg/ml. The results showed that most of the derivatives have good cytotoxic activity but poor antiviral activity [12].

Vinayak et al. synthesized novel phenoxy acetamide derivatives (fig. 11) where amide nitrogen is substituted with the heterocyclic ring and evaluated these compounds for MTT (Tetrazolium dye) assay using various cell lines such as HepG2 (human liver cancer cell line) and Caco-2 (human colonic epithelial cell line). Among the synthesized compounds, derivative bearing fluoro substitution at 2nd position was found to be highly active on Caco-2 cell line with IC₅₀ of 1.8 μmol whereas other compounds showed less cytotoxicity on all the three cell lines as compared with the standard drug 5-fluorouracil [13].

Antibacterial and antifungal activity

A series of novel is substituted pyrazolo phenoxy acetic acid derivatives (fig. 13) were synthesized and evaluated them for antitubercular activity using Middlebrook 7H9 agar medium against H₃7Ra Strain of M. tuberculosis. Most of the synthesized derivatives displayed significant antitubercular activity [15].

A series of azomethine derivatives of phenoxy acetic acid (fig. 16) were synthesized by treating 2-formylphenoxyacetic acid with aromatic amines such as 2,4-dimethylpyridine and 2,3-dichloro aniline. These compounds were assayed by the disc diffusion method for antibacterial activity using several bacteria such as Staphylococcus aureus and Escherichia coli.

Among the derivatives tested, p-aminoacetanilide, 2, 3-dichloro aniline and p-toluidine containing derivatives exhibited good antibacterial activity, similar to that of standard drug ciprofloxacin [18].
Kaplancikli et al. synthesized a series of phenoxy acetamide derivatives substituted with carbazole moiety (Fig. 17) by treating phenoxy acetamide with various ring-substituted phenols. These compounds were screened for their antibacterial and antifungal activities against *Micrococcus luteus, Bacillus subtilis, Staphylococcus aureus, Listeria monocytogenes* and *Candida albicans*.

These compounds showed notable antimicrobial activity against the tested organisms. The compounds were also studied for their cytotoxic effects using MTT assay, where quinoline bearing derivative had the poor cytotoxic activity against NIH 3T3 (the murine fibroblast) cells [19].

**MDR (Multidrug resistance) reversing activity**

P-glycoprotein (Pgp) can actively transport anticancer agents out of the cancer cells and decrease their intracellular accumulation. Pgp has emerged as a promising target for cancer therapy and in the development reversal agents to overcome Pgp-mediated MDR. Adamantyl based Pgp inhibitor (Fig. 18) is highly potent which has phenoxy acetamide unit in its structure.

Lee *et al.* synthesized a series of phenoxy-N-phenylacetamide derivatives (Fig. 19) and tested for their ability to inhibit Pgp, using a Pgp over-expressing breast cancer cell lines (MCF-7/ADR). Derivative with methyl furyl substitution showed a 3-fold increased inhibition compared with verapamil, a potent Pgp inhibitor, whereas rest of the compounds exhibited moderate activity [20].

**Cholecystokinin-B receptor antagonistic activity**

Takeda *et al.* synthesized various phenoxy acetamide derivatives (Fig. 20) and evaluated them for their cholecystokinin-B receptor antagonistic activity. Most of the derivatives exhibited good affinity for these receptors, and structure-activity relationship showed the importance of the optimal size of the N-alkyl chains for the inhibitory activity. Derivatives bearing cyclohexyl ring or methyl pentyl side chain demonstrated the highest activity [21].

**Antihypertensive activity**

Various phenoxy acetamide derivatives containing pyridazine heterocyclic ring at para position (Fig. 21) were synthesized, and their antihypertensive effects were evaluated using *in vitro* method using standard reference drug hydralazine. Derivative having pyrrolidine ring on amide nitrogen showed moderate antihypertensive activity whereas introduction of other heterocyclic rings such as piperidine, morpholine, phenyl hydrazine and aniline significantly reduced the antihypertensive activity [22].

**Antioxidant activity**

Prashanth *et al.* synthesized a series of (4-benzoyl-phenoxy)-acetic acid derivatives (Fig. 22) and tested *in vitro* antioxidant effects by using 1,1-diphenyl-1-picrylhydrazyl (DPPH), nitric oxide (NO) and hydrogen peroxide (H$_2$O$_2$) radical scavenging assays. Among the synthesized compounds, unsubstituted derivative and derivative possessing chlorine atom on benzoyl ring showed good radical scavenging activity in all the three methods compared to the standard drug ascorbic acid. Derivative containing methoxy substituent on benzoyl ring showed good antioxidant activity only in hydrogen peroxide method and rest of the compounds showed moderate to poor radical scavenging activity [23].
A novel series of phenoxy acids containing chalcone scaffold (fig. 23) were synthesized and screened for their antioxidant activity using DPPH free radical scavenging assay. All the synthesized compounds were found to display good activity in this assay indicating that these structures are potential antioxidant agents [24].

Salicylamidoacetic acid hydrazide (6) was found to have good analgesic and antiinflammatory properties and fewer side effects. A series of phenoxy hydrazide derivatives (fig. 25) were synthesized to obtain safe and effective agents [28].

**Fig. 23**

**Miscellaneous activities**

Phenoxy acid containing chemical compounds are versatile for their herbicidal properties. Several selective herbicides such as 2, 4-Dichloro phenoxy acetic acid (3), Clomeprop (4) and Difenopenten (5) contain phenoxy acid moiety as a part of their structure [25].

A series of phenoxy acetamide derivatives possessing substituted heterocyclic moiety (fig. 23) were synthesized using tetrabutylammonium bromide as phase transfer catalyst from corresponding o-fluorophenol and evaluated for their herbicidal activities. The preliminary herbicidal tests showed that the synthesized compounds have good inhibitory activities against root and stalk of dicotyledon plants [26].

**Fig. 24**

A series of phenoxy isobutyric acid derivatives (fig. 24) in the form of choline salt were synthesized which were observed to have photostability and better reactivity [27].

A novel series of phenoxy containing amide linker and various substituted heterocyclic rings such as pyrazole (fig. 28) were synthesized and evaluated for their PPAR-γ inhibitory activity. Most of the derivatives displayed good activity and derivative possessing methyl functionality at 3rd position displayed the highest activity in the in vitro and in vivo models [33].

**Fig. 28**

**CONFLICT OF INTERESTS**

Declared none

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