

Histamine Receptors

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Dr. Iwan de Esch is an assistant professor and Prof. Rob Leurs is full professor and head of the Division of Medicinal Chemistry of the Leiden/Amsterdam Center of Drug Research (LACDR), VU University Amsterdam, The Netherlands. Since the seventies, histamine receptor research has been one of the traditional themes of the division. Molecular understanding of ligand-receptor interaction is obtained by combining pharmacology (signal transduction, proliferation), molecular biology, receptor modelling and the synthesis and identification of new ligands.

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Introduction

Histamine is one of the aminergic neurotransmitters and plays an important role in the regulation of several (patho)physiological processes. In the mammalian brain histamine is synthesised in restricted populations of neurons that are located in the tuberomammillary nucleus of the posterior hypothalamus.1 These neurons project diffusely to most cerebral areas and have been implicated in several brain functions (e.g. sleep/ wakefulness, hormonal secretion, cardiovascular control, thermoregulation, food intake, and memory formation).2 In peripheral tissues, histamine is stored in mast cells, eosinophils, basophils, enterochromaffin cells and probably also in some specific neurons. Mast cell histamine plays an important role in the pathogenesis of various allergic conditions. After mast cell degranulation, release of histamine leads to various well-known symptoms of allergic conditions in the skin and the airway system. In 1937, Bovet and Staub discovered compounds that antagonise the effect of histamine on these allergic reactions.3 Ever since, there has been intense research devoted towards finding novel ligands with (anti-) histaminergic activity. This research field has been fuelled by the consecutive discovery of four unique histamine receptor subtypes. Every receptor subtype has a very distinct (patho)physiological role and all of them belong to the superfamily of G-protein-coupled receptors (GPCRs). The histamine H₁ and H₂ receptor subtypes have proven to be excellent drug targets. Ligands for the histamine H₂ receptor subtype are currently entering clinical studies and the recently discovered histamine H₄ receptor subtype is subject of intense preclinical research.

The Histamine H₁ Receptor

Until the seventies, histamine research focused on the role of histamine in allergic diseases. This resulted in the development of several potent 'antihistamines' (e.g. mepyramine, see figure 1), which were useful in inhibiting pronounced symptoms of allergic conditions.4 However, the first generation of H₁ receptor antagonists that were developed for treating allergies revealed distinct side effects such as sedation. This particular physiological effect of the ligands was eliminated by structural modifications that prevent blood-brain-barrier penetration of the drugs. The first generation as well as the more recently developed antihistamines (originally termed antagonists but later reclassified as inverse agonists5) are clinically very successful and are widely available drugs. In 1991, the cDNA encoding a bovine H₄ receptor protein was cloned after an expression cloning strategy in Xenopus oocytes.6 The deduced amino acid sequence revealed a 491 amino acid protein of 56 kDa. Using the cDNA sequence encoding the bovine H₁ receptor, the cDNA sequences and intronless genes encoding the rat, 3,7 guinea-pig, 8,9 human 10,11 and mouse¹² H₄ receptor proteins were cloned soon thereafter. These receptor proteins are slightly different in length, highly homologous and do not show major pharmacological differences. Analysis of the 5'-flanking region of the human, rat and guineapig gene^{7,8,10} resulted in the identification of several DNA-binding motifs, including potential glucocorticoid responsive elements. The human H₄ receptor gene resides on chromosome 3.13 The H₄ receptor belongs to the large family of GPCRs. The receptor is associated with the phospholipase C-catalysed formation of inositol 1,4,5-triphosphate (IP₃) and 1,2-diaylglycerol (DAG). Histamine induces production of inositol phosphates in several tissues (including brain, airway, intestinal and vascular smooth $\text{muscle}^{24})$ via $\mathbf{G}_{\alpha \mathbf{a}}$ protein activation. 14 In other tissues, activation of $\mathrm{H_{\scriptscriptstyle{4}}}$ receptors can also stimulate adenylyl cyclase and formation of cAMP. Not all details of this signalling pathway are understood, for example questions remain about the G protein and the involvement of Ca²⁺. ¹⁵ In any case, alternative signalling pathways can be mediated by the histamine H₁ receptor. Recent results indicate that the functional heterogeneity can be liganddirected.16

The histamine $\rm H_1$ receptor is a well established drug target and has been thoroughly studied for decades. Nevertheless, $\rm H_1$ receptor research continues to flourish as many new techniques and approaches are being developed by using this receptor as an archetypal GPCR target. Most notably, in the last few years, the histamine $\rm H_1$ receptor has been the subject of various molecular biology studies (e.g., large-scale overproduction and GPCR

binding site elucidation 18), biophysical approaches (such as solid-state NMR 19) and investigations towards the general activation mechanisms of GPCRs. 20,21

Ligands for H₁ Receptors

Modification of the imidazole moiety of histamine has been the most successful approach for obtaining selective H, agonists (figure 1). The presence of the tautomeric N^{π} - N^{τ} system of the imidazole ring is not obligatory, as reflected by the selective H₄ agonists 2-pyridylethylamine and 2-thiazolylethylamine. Substitution of the imidazole ring at the 2-position leads to relatively selective H_a agonists. For example, 2-(meta-halogenated) phenylhistamines are relatively potent H, receptor agonists at the guinea-pig ileum;22 however, these compounds act as partial agonists in other systems.²³ A wide array of potent and selective H₄ antagonists are available.⁴ Compounds such as mepyramine (also called pyrilamine) and triprolidine are highly potent H, antagonists and very useful tools for pharmacological investigations. [3H]-mepyramine is, for example, successfully used as an H₁ receptor radioligand.²⁴ These so-called classical 'antihistamines' easily penetrate the brain and are therefore also useful in in vivo CNS studies.2 Elimination of the blood-brainbarrier passage by some minor structural modifications has resulted in many new, non-sedating H₁ antagonists (e.g., cetirizine, astemizole, fexofenadine and loratidine).4

Figure 1 | Chemical structures of selected H, receptor ligands

(Bold text denotes compounds available from Tocris at time of publication.)

Figure 2 | Chemical structures of selected H₂ receptor ligands

(Bold text denotes compounds available from Tocris at time of publication.)

The Histamine H₂ Receptor

The observation that the classical 'antihistamines' (i.e. H₁ receptor inverse agonists) cannot antagonise all histamineinduced effects (e.g. at the stomach and the heart), led Ash and Schild in 1966 to propose two distinct subtypes of histamine receptors: H₁ and H₂.²⁵ This hypothesis became generally accepted when Black et al26 succeeded in the synthesis of a series of new compounds (e.g. burimamide, cimetidine), which were able to block the effects of histamine on the stomach and the heart. These H₂ receptor antagonists proved to be very useful in the therapy of gastric ulcers. Gantz et al27 were the first to clone a cDNA encoding a 359 amino acid H₂ receptor. Using degenerate primers based on the known sequence similarity of various GPCRs, the H₂ receptor sequence was obtained from canine gastric parietal cDNA by PCR. Soon thereafter, the intronless genes encoding the rat,28 human,29 guinea pig30 and mouse³¹ H₂ receptor were cloned by means of homology screening. Identification of the promotor region of the human H₂ receptor gene revealed the existence of regulatory transcription sites and regions displaying stimulatory and inhibitory effects on gene expression monitored in a luciferase assay.32 Studies have indicated that the human H2 receptor gene resides on chromosome 5.30 Interestingly, several polymorphisms have been found in the human H_2 receptor gene³³ and one of the mutations has been linked to schizophrenia.34 The histamine H₂ receptor is coupled to the adenylate cyclase system in a variety of tissues (e.g. brain, stomach, heart, gastric mucosa, lung).24 Moreover, cell lines transfected with the cloned H₂ receptor genes showed an H₂ receptor-mediated increase of cAMP. 35,36,37 In addition, alternative signalling pathways for the H2 receptor have been identified. In differentiated HL-60 cells and CHO or HEPA cells transfected with the H₂ receptor cDNA, an H₂ receptor-mediated increase of the intracellular Ca2+ concentration and/or IP3 levels was observed.^{38,39} Moreover, in CHO cells expressing the rat H₂ receptor, activation of the H₂ receptor resulted in an inhibition of the release of arachidonic acid induced by either constitutive purinergic receptors or a Ca²⁺-ionophore,³⁶ as well as an increase in cAMP. These new signal transduction pathways are both regulated via unknown, cAMP-independent pathways.

Ligands for H₂ Receptors

The first selective $\rm H_2$ receptor agonist was dimaprit (figure 2). This compound is almost as active as histamine at the $\rm H_2$ receptor but hardly displays any $\rm H_1$ receptor activity. Much later, it was shown that dimaprit is also a moderate $\rm H_3$ receptor antagonist and a moderate $\rm H_4$ receptor agonist. Amthamine can be considered a rigid dimaprit analogue. This compound combines a high $\rm H_2$ receptor selectivity with a potency which is slightly higher compared to histamine, both *in vitro* and *in vivo*. An $\rm H_2$ receptor agonist that is more potent than histamine is the guanidine derivative impromidine. This ligand actually combines a rather high $\rm H_2$ receptor affinity with a reduced efficacy. Impromidine also shows moderate $\rm H_1$ - and potent $\rm H_3$ -receptor antagonistic activity. So well as potent $\rm H_4$ receptor partial agonistic activity.

The finding that N α -guanylhistamine acts as a partial H $_2$ agonist in a gastric acid secretion test did lead to the development of the relatively weak H $_2$ antagonist burimamide. Years later, it was shown that burimamide is also an H $_3$ and H $_4$ receptor partial agonist. Nevertheless, burimamide was a good lead for the development of selective and clinically useful H $_2$ receptor antagonists, such as cimetidine. The 4-methylimidazole moiety of cimetidine can easily be replaced by other heterocyclic groups (figure 2). Replacement by a substituted furan- (e.g. ranitidine) or thiazole ring (e.g. tiotidine and famotidine) leads to compounds that are usually more potent at the H $_2$ receptor than

(R)-(-)-α-Methylhistamine, High Affinity H₂ Agonist

(R)-(-)- α -Methylhistamine

Cat. No. 0569

This potent and high affinity $\rm H_3$ agonist displays >200-fold selectivity over $\rm H_4$ receptors. The compound inhibits $\rm H_3$ receptor-mediated histamine synthesis and release in the CNS and stimulates $\rm H_4$ receptor-mediated eosinophil shape change (EC $_{50}$ = 66 nM).

Oishi et al. (1989) Effects of histamine H_3 -agonist (R)- α -methylhistamine and the antagonist thioperamide on histamine modulation in the mouse and rat brain. J.Neurochem. 52 1388. Hew et al. (1990) Characterization of histamine- H_3 receptors in guinea pig illeum with H_3 -selective ligands. Br.J.Pharmacol. 101 621. Schwartz et al. (1990) A third histamine receptor subtype - characterization, localization and functions of the H_3 -receptor. Agents Actions 30 13. Buckland et al. (2003) Histamine induces cytoskeletal changes in human eosinophils via the H_4 receptor. Br.J.Pharmacol. 140 1117.

cimetidine. Moreover, the replacement of the imidazole moiety also eliminates the undesired inhibition of cytochrome P450. The potent tritiated $\rm H_2$ antagonist tiotidine and [125 I]-iodinated $\rm H_2$ antagonist iodoaminopotentidine are successfully used as radioligands for the $\rm H_2$ receptor. The newly developed brain-penetrating $\rm H_2$ antagonist zolantidine is an important tool for *in vivo* CNS studies. The $\rm H_2$ receptor was reported to be spontaneously active in transfected CHO cells. Based on this concept, many $\rm H_2$ antagonists were reclassified; cimetidine, ranitidine and famotidine are in fact inverse agonists, whereas burimamide acts in this model system as a neutral antagonist.

The Histamine H₃ Receptor

In the eighties, another physiological role of histamine became apparent, namely its role as a neurotransmitter. In 1983, Arrang and co-workers discovered that the inhibitory effect of histamine on its own release and synthesis was not mediated by the known H₁ and H₂ receptor subtypes as no correlation with either the H₁ or the H₂ receptor activity of known histaminergic compounds was observed.⁴⁶ Soon after, the H_3 receptor agonist (R)-(-)- α methylhistamine and the antagonist thioperamide (see figure 3) were developed.51 It was confirmed that this receptor subtype indeed regulates the release and synthesis of histamine and in addition has a regulatory role in the release of other neurotransmitters, such as serotonin, noradrenalin and dopamine.52 Next to high expression in certain regions of the CNS (for example the basal ganglia, hippocampus and cortical areas, i.e. the parts of the brain that are associated with cognition) the H₃ receptor is present in the peripheral nervous system, e.g. in the gastrointestinal tract, the airways and the cardiovascular system. 53,54 Initial efforts to identify the H₃ receptor gene, using the anticipated homology with the identified H₁ and H₂ receptor gene all failed. Eventually, the human H₃ receptor cDNA was cloned by Lovenberg and co-workers in 1999.55 In search of novel GPCRs through homology searching of expressed sequence tag databases, a receptor with high similarity to the M₂ muscarinic acetylcholine receptor was identified. Expression of the gene and full characterisation established this protein as the histamine H_a receptor. The cloning of the H₃ receptor of other species, including rat, 56 guinea pig 57 and mouse 58 soon followed and it was revealed that major $\rm H_3$ receptor species differences exist. The human $\rm H_3$ receptor gene was assigned to the telomeric region of the q arm of chromosome 20 and contains three exons that are interrupted by two introns. 59 The $\rm H_3$ receptor gene can undergo extensive alternative splicing, resulting in many $\rm H_3$ receptor isoforms that have different signalling properties and expression profiles. 59,60,61 It was shown that the $\rm H_3$ receptor displays particularly high constitutive activity, again leading to a reclassification of existing ligands into agonists, neutral antagonists and inverse agonists. 62,63 Gbahou and co-workers revealed that the compound proxyfan (figure 3) is a protean agonist both *in vitro* and *in vivo*, meaning that this remarkable compound behaves as agonist, neutral antagonist or inverse agonist, depending on the signalling pathway studied. 64

The H₃ receptor signals through G_{1/0} proteins as was shown by the pertussis toxin sensitive stimulation of [35S]-GTPγS binding in rat cortical membranes.65 Through negative coupling to adenylyl cyclase, stimulation of the H3 receptor results in lower levels of cAMP, thereby reducing downstream signalling events such as CREB-dependent gene transcription.56 Alternative signalling pathways may be activated by the G_{1/0} proteins, including mitogenactivated protein kinase (MAPK)60 and phosphatidylinositol 3-kinase (PI3K) pathways. G_{1/0} protein activation can also lead to the activation of phospholipase A2 (PLA2) to induce the release of arachidonic acid, the lowering of intracellular Ca2+ levels through voltage-gated ion channels⁶⁷ and the inhibition of the Na⁺/H⁺ exchanger (NHE).68 With the recent progress that has been made in the characterisation of the H₃ receptor (as outlined above), many pharmaceutical companies have very active H₃ receptor drug development programs. As a result, several H3 receptor ligands have entered clinical studies for a plethora of applications, including obesity, narcolepsy, dementia and migraine (among others).69,70

Ligands for H₃ Receptors

At the H₃ receptor, histamine itself is a highly active agonist. Modification of the endogenous ligand by mono- or dimethylation of the amino function results in compounds that are more active

Thioperamide, a Brain-penetrant H₂ and H₄ Ligand

Thioperamide

Cat. No. 0644

Thioperamide, the potent histamine $\rm H_3$ and $\rm H_4$ ligand, displays $\rm K_i$ values of 25 and 27 nM at recombinant $\rm H_3$ and $\rm H_4$ receptors respectively. The compound acts as an antagonist at $\rm H_3$ receptors and displays inverse agonist activity at $\rm H_4$ receptors. It freely crosses the blood-brain barrier.

Hew *et al.* (1990) Characterisation of histamine $\rm H_3$ receptors in guinea pig ileum with $\rm H_3$ selective ligands. *Br.J.Pharmacol.* **101** 621. Liu *et al.* (2001) Cloning and pharmacological characterization of a fourth histamine receptor ($\rm H_4$) expressed in bone marrow. *Mol. Pharmacol.* **59** 420. Ling *et al.* (2004) Histamine $\rm H_4$ receptor mediates eosinophil chemotaxis with cell shape change and adhesion molecule upregulation. *Br.J.Pharmacol.* **142** 161. **Gbahou** *et al.* (2006) Compared pharmacology of human $\rm H_3$ and $\rm H_4$ receptors: structure-activity relationships of histamine derivatives. *Br.J.Pharmacol.* **147** 744.

and selective for the H_3 receptor. Methylation of the α -carbon atom of the ethylamine sidechain drastically increases the potency at the H₃ receptor. This increased activity resides completely in the R-isomer; the corresponding S-isomer is approximately 100-fold less potent. Since the methylation leads to highly reduced activity at both H₁ and H₂ receptors, (R)-(-)-αmethylhistamine (figure 3) was initially considered a selective agonist at the H_a receptor. However, when the H₄ receptor was discovered it was shown that (R)-(-)- α -methylhistamine also has considerable affinity for this histamine receptor subtype.42 Nevertheless, in combination with its less active S-isomer, (R)-(-) $-\alpha$ -methylhistamine has proven to be highly useful for the pharmacological characterisation of H₃ receptor-mediated effects. 41 Tritiated forms of N α -methylhistamine and (R)-(-)- α methylhistamine are available as radiolabelled agonists for the $\mathrm{H_{3}}$ receptor. For potent $\mathrm{H_{3}}$ agonism, the amine function of histamine can be replaced by an isothiourea group, as in imetit. This compound is also very active in vitro and in vivo, 71,72,73 as is (R)-(-)- α -methylhistamine. The amine function can also be incorporated in ring structures. For example, immepip is a potent H₃ receptor agonist that is effective in vitro and in vivo.⁷⁴ Although the described first generation H₃ agonists were intensively used as reference ligands to study the H₃ receptor, all of them proved to also have considerable activity at the ${\rm H_4}$ receptor. Therefore, a new generation of potent and selective H₃ receptor agonists has been developed, most notably immethridine⁷⁵ (pEC₅₀ = 9.74; displays 300-fold selectivity over the H_4 receptor) and

Imetit, High Affinity H./H. Agonist

Imetit

Cat. No. 0729

Imetit is an extremely high affinity, potent agonist at $\rm H_3$ and $\rm H_4$ receptors ($\rm K_i$ values are 0.3 and 2.7 nM respectively). The agonist induces shape change in eosinophils *in vitro* with an EC₅₀ of 25 nM and is centrally active following systemic administration *in vivo*.

Garbarg et al. (1992) S-[2-(4-Imidazolyl)ethyl]isothiourea, a highly specific and potent histamine $\rm H_3$ receptor agonist. J.Pharmacol.Exp.Ther. 263 304. Farzin and Attarzadeh (2000) Influence of different histamine receptor agonists and antagonists on apomorphine-induced licking behavior in rats. Eur.J.Pharmacol. 404 169. Ling et al. (2004) Histamine $\rm H_4$ receptor mediates eosinophil chemotaxis with cell shape change and adhesion molecule upregulation. Br.J.Pharmacol. 142 161.

methimepip. 76 These latter compounds are devoid of high $\rm H_4$ receptor activity.

As with the first generation $\rm H_3$ receptor agonists, the first generation $\rm H_3$ receptor antagonists (all of them possessing an imidazole heterocycle) turned out to have affinity for the $\rm H_4$ receptor. The first potent $\rm H_3$ receptor antagonist (later reclassified as an inverse agonist) that was devoid of $\rm H_4$ receptor and $\rm H_2$

Figure 3 | Chemical structures of selected H₂ receptor ligands

receptor activity was thioperamide (figure 3).51 This compound has been used in many H_a receptor studies as a reference ligand and is active in vitro and in vivo (the compound is able to penetrate the CNS).77 However, thioperamide displays some 5-HT₃ receptor antagonism⁷⁸ and is an inverse agonist at the H₄ receptor. The remarkable H₃ receptor species difference can be demonstrated with thioperamide as the compound has a 10-fold higher affinity for the rat H₃ receptor than for the human H₃ receptor.56 Based on the H₃ receptor agonist imetit (vide ante), the highly potent H_a receptor inverse agonist clobenpropit was developed (pA₂ = 9.9).⁷³ This compound has some 5-HT₃ receptor activity⁷⁸ and displays partial agonist activity at H₄ receptors. In addition, radioligands for the H_3 receptor have been described (e.g. $[^{125}I]$ -iodophenpropit and $[^{125}I]$ -iodoproxyfan). 79,80 The moderately active H_2 receptor antagonist burimamide (pA₂ = 5.1, figure 2) also has good affinity for the H_a (pK_i = 7.9) and the H_a $(pK_1 = 7.4)$ receptor.⁴² Impentamine is a potent histamine H₂ receptor inverse agonist (pA $_{2}$ = 8.4). Like burimamide, this compound can act as a partial agonist in SK-N-MC cells expressing human H_a receptors. It has also been shown that small structural modifications of impentamine, i.e. alkylation of the primairy amine moiety of impentamine with e.g. methyl-, isopropyl- and p-chlorobenzyl- groups results in ligands that cover the complete spectrum of functional activity, i.e. agonism, neutral antagonism and inverse agonism. 63,81 The compound VUF 5681 was reported as a neutral H₂ antagonist (or 'silent' antagonist),82 findings that were recently confirmed in studies investigating constitutive activity using rat brain cortex.83 It has been shown that the functional activity of proxyfan depends on the system used, ranging from full agonist to inverse agonist. Thus, proxyfan can be classified as a protean agonist, both in vitro and in vivo. 64,84

The first non-imidazole H₃ receptor ligand was reported by Ganellin in 1998.85 This elegant medicinal chemistry work did lead to the potent compound UCL 2190.86 Following the cloning of the H₃ receptor in 1999, several pharmaceutical companies entered the H₂ research field and further explored this structural class. An example of these new ligands is JNJ 5207852, a compound that contains some typical structural features for H₃ receptor ligands, e.g. piperidinyl propyloxy side chain. This neutral antagonist is active in several models for cognition87,88 but does

Clobenpropit, Highly Potent H₂ Antagonist

Clobenpropit

Cat. No. 0752

An extremely potent antagonist/inverse agonist at H₃ receptors (pA₂ = 9.93), clobenpropit also displays partial agonist activity at H₄ receptors. The ligand induces eosinophil shape change with an EC_{50} of 3 nM.

Van der Goot et al. (1992) Isothiourea analogues of histamine as potent agonists or antagonists of the histamine H, receptor. Eur.J.Med.Chem. 27 511. Yokoyama et al. (1994) Clobenpropit (VUF-9153), a new histamine H₃ receptor antagonist, inhibits electrically induced convulsions in mice. Eur.J.Pharmacol. 260 23. Liu et al. (2001) Cloning and pharmacological characterization of a fourth histamine receptor (H_a) expressed in bone marrow. Mol. Pharmacol. 59 420. **Buckland** et al. (2003) Histamine induces cytoskeletal changes in human eosinophils via the H₄ receptor. Br.J.Pharmacol. 140 1117.

VUF 5681, a Novel H₂ Antagonist

VUF 5681

Cat. No. 2493

VUF 5681 is a potent histamine H₂ receptor silent antagonist

Kithunnadai et al. (2004) Identification of 4-(1H-imidazol-4(5)-vlmethyl) pyridine (immethridine) as a novel, potent, and highly selective histamine H receptor agonist. J.Med.Chem. 47 2414. Leurs et al. (2005) The histamine H. receptor: from gene cloning to H3 receptor drugs. Nat.Rev.Drug Discov. 4 107. Moreno-Delgado et al. (2006) Constitutive activity of H₂ autoreceptors modulates histamine synthesis in rat brain through the cAMP/PKA pathway. Neuropharmacology 51 517.

not act as an appetite suppressant and has no effect on food intake.89 Other compounds, such as Abbott's A 423579 seem to have more efficacy in obesity models, while lacking clear cognitive effects.90 At the time of writing the differences in efficacy for distinct clinical applications of the different classes of H_a ligands is not understood and subject of intense research. 70,91 Interestingly, GSK 189254 has been in trials for three different diseases: neuropathic pain, narcolepsy and dementia.92

The Histamine H₁ Receptor

Immediately following the cloning of the H₃ receptor gene, several groups identified the homologous H₄ receptor sequence in the human genome databases. $^{93-97}$ Indeed, the H_4 receptor has high sequence identity with the H₂ receptor (31% at the protein level, 54% in the transmembrane domains). The H₂ and H₄ receptor are also similar in gene structure. The human H, receptor gene is located on chromosome 18q11.2 in a single copy per haploid genome. The gene spans more than 21 kbp and contains three exons that are interrupted by two large introns.98 To date, two H_a receptor isoforms have been identified. Cloning of the genes that encode the mouse, rat, guinea-pig and pig H₄ receptors reveal only limited sequence homology with the human H₄ receptor. The H₄ receptor is mainly expressed in bone marrow and peripheral leukocytes, and mRNAs of the human H, receptor are detected in e.g. mast cells, dentritic cells, spleen and eosinophils.93-97 The receptor has a pronounced effect on the chemotaxis of several cell types that are associated with immune and inflammatory responses. The H_4 receptor is mainly coupled to $G_{i/o}$ proteins, thereby leading to a decrease in the production of cAMP and subsequent downsteam effects such as regulation of cAMP responsive element-binding protein (CREB) gene transcription. Furthermore, $H_{\scriptscriptstyle \Delta}$ receptor stimulation affects the pertussin-toxin-sensitive activation of mitogen-activitated protein (MAP) kinase pathways. Studying the increased levels of [35S] GTPyS levels in H₄ transfected cells, it has been shown that the H₄ receptor is constitutively active. The $G\beta\gamma$ subunits of the $G_{i/o}$ proteins activate phospholipase C, and thereby increase the Ca2+ concentrations. In mast cell and eosinophils, this Ca2+ response can be linked to cellular chemotaxis.

Considering the physiological role of the H_a receptor, several applications are under preclinical investigation, 99,100 including allergy and asthma,101 as well as chronic inflammations such as

Figure 4 | The first selective H4 receptor ligands reported in scientific literature

(Bold text denotes compounds available from Tocris at time of publication.)

JNJ 10191584, H₄-Selective Antagonist

JNJ 10191584 is a highly selective histamine $\rm H_4$ receptor silent antagonist. It binds with high affinity to the human $\rm H_4$ receptor ($\rm K_i$ = 26 nM) and is > 540-fold selective over the $\rm H_3$ receptor ($\rm K_i$ = 14.1 $\rm \mu M$). In vitro the antagonist inhibits mast cell and eosinophil chemotaxis with IC $_{50}$ values of 138 and 530 nM respectively. The antagonist is orally active *in vivo*.

Terzioglu et al. (2004) Synthesis and structure-activity relationships of indole and benzimidazole piperazines as histamine $\rm H_4$ receptor antagonists. Bioorg. Med. Chem.Lett. **14** 5251. **Venable** et al. (2005) Preparation and biological evaluation of indole, benzimidazole, and thienopyrrole piperazine carboxamides: potent human histamine $\rm H_4$ antagonists. J.Med.Chem. **48** 8289. **Varga** et al. (2005) Inhibitory effects of histamine H4 receptor antagonists on experimental colitis in the rat. Eur.J.Pharmacol. **522** 130.

inflammatory bowel disease (IBD) 102 and rheumatoid arthritis. 103 The H $_4$ receptor is also being associated with pruritus (itch) 104,105 and is involved in the progression of colon cancer. 106

Ligands for H₄ Receptors

As was described above, most imidazole-containing, first generation $\rm H_3$ receptor ligands have considerable affinity for the $\rm H_4$ receptor as well. However, selective $\rm H_4$ receptor ligands have been described. 4-Methylhistamine is a potent $\rm H_4$ agonist while displaying more than a 100-fold selectivity over the other histamine receptor subtypes, including the $\rm H_2$ receptor for which this ligand was originally developed. A slightly different and complimentary profile was reported for VUF 8430. This compound has a high $\rm H_4$ receptor activity and affinity, minimal affinity for the $\rm H_1$ and $\rm H_2$ receptor and a 33-fold lower affinity for the $\rm H_3$ receptor.

Potent and selective $\rm H_4$ receptor antagonists are also emerging. The first reported neutral antagonist is JNJ 7777120, a compound that is frequently being used as $\rm H_4$ receptor reference ligand. Currently, [³H]-JNJ 7777120 and [³H]-histamine are used as $\rm H_4$ receptor radioligands. Unfortunately, JNJ 7777120 has a poor stability in human and rat liver microsomes and a limited half life of about two hours. The benzimidazole derivative JNJ 10191584 (VUF 6002) is also a neutral $\rm H_4$ antagonist. This compound is orally active *in vivo* and has an improved liver microsome stability but still a limited half life. 102,109 More recently, 2-arylbenzimidazoles

have been described as ligands with low nanomolar affinity for the $\rm H_4$ receptor. ^110 Considering the number of $\rm H_4$ receptor-related patent applications that have recently been disclosed (as reviewed elsewhere ^100), it can be anticipated that many new $\rm H_4$ receptor ligands will be described in scientific literature in the near future.

Conclusions

The histamine receptor research field continues to blossom as both members of industry and academia find this family of receptors very rewarding, both scientifically and commercially. For more than seventy years, considerable efforts have been devoted to finding new ways to modulate the different physiological processes that are mediated by histamine. The consecutive discovery of new histamine receptor subtypes (distinction between H₁ and H₂ receptors in 1966, discovery of the H₃ receptor in 1983 and the H₄ receptor in 2000, note the interval of seventeen years) provides a complete and exciting research platform. The histamine receptor family also seems to hold the promise of GPCRs as excellent drug targets with two receptor subtypes (H_a and H_a) addressed by blockbuster drugs, the third subtype (H₂) leading to frantic clinical studies and the latest addition to the family (H₄) leading to very interesting preclinical data. It can therefore be anticipated that histamine receptor research will continue to thrive in the years to come.

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Histamine Receptor Compounds Available from Tocris

Catalog No.	Product Name	Primary Function	
H, Receptors			
Agonists			
2478	2-Pyridylethylamine	H ₁ agonist	
Inverse Agonists			
0660	Mepyramine	Selective H ₁ inverse agonist	
Antagonists			
3489	<u>Astemizole</u>	Orally active, potent $\rm H_{1}$ antagonist; also $\rm K_{v}11.1$ (hERG) channel blocker	
2577	Cetirizine	Selective H ₁ antagonist	
1453	Clemastine	H ₁ antagonist	
5371	Clemizole	H ₁ antagonist; also TRPC5 blocker	
5958	<u>Desloratadine</u>	High affinity H ₁ antagonist; anti-inflammatory	
1425	(S)-(+)-Dimethindene	${\rm H_1}$ antagonist; also ${\rm M_2}$ muscarinic antagonist	
3072	<u>Diphenhydramine</u>	H ₁ antagonist	
0508	<u>Doxepin</u>	Highly potent $\rm H_1$ antagonist; also binds to $\rm H_4$ receptor	
2429	<u>Fexofenadine</u>	${\rm H_1}$ receptor antagonist; non-sedating antiallergic agent	
0784	Ketotifen	H ₁ antagonist	
1944	Loratidine	Peripheral H ₁ antagonist; anti-allergic agent	
4245	Meclizine	${\rm H_1}$ antagonist; also human pregnane X receptor agonist	
2018	<u>Mirtazepine</u>	Potent $\rm H_1$ antagonist; also 5-HT $_{\rm 2}$, 5-HT $_{\rm 3}$ and $\alpha_{\rm 2}$ -antagonist; antidepressant	
4241	<u>Olopatadine</u>	H ₁ antagonist; ocular anti-allergic agent	
3948	<u>Terfenadine</u>	$\rm H_{1}$ antagonist; also $\rm K_{\rm v}11.1$ (hERG) and $\rm K_{\rm ir}6$ (KATP) channel blocker	
0662	<u>trans-Triprolidine</u>	Highly potent H ₁ antagonist	
3996	Zotepine	$\rm H_1$ antagonist; also 5-HT $_{\rm 2A}$ and $\rm D_2$ antagonist	
H ₂ Receptors			
Agonists			
0668	<u>Amthamine</u>	Highly selective standard H ₂ agonist	
0506	Dimaprit	Standard H ₂ selective agonist	
Antagonists			
0721	Aminopotentidine	H ₂ antagonist	
0902	Cimetidine	H ₂ antagonist; also I ₁ agonist	
0833	<u>ICI 162,846</u>	Potent H ₂ antagonist; active <i>in vivo</i>	
1967	Ranitidine	Selective H ₂ antagonist	
0826	Tiotidine	Potent and selective H ₂ antagonist	
1070	Zolantidine	Potent, centrally active H ₂ antagonist	
H ₃ Receptors			
Agonists			
2315	Immethridine	Potent H ₃ agonist; highly selective over H ₄	
0573	<u>N</u> ^a _Methylhistamine	Non-selective H ₃ agonist	
0569	(R)-(-)-α-Methylhistamine	Potent standard H ₃ agonist	
0572	(S)-(+)-α-Methylhistamine	H ₃ agonist; less active enantiomer	
2477	<u>Proxyfan</u>	High affinity H ₃ ligand	
Inverse Agonists			
3743	BF 2649	Potent and selective H ₃ inverse agonist	
Antagonists			
4697	<u>A 331440</u>	Selective H ₃ antagonist	
2211	Carcinine	Highly selective H ₃ antagonist	
2419	GT 2016	Selective H ₃ antagonist	
1858	<u>Impentamine</u>	Selective H ₃ antagonist	
0779	lodophenpropit	Potent and selective H ₃ antagonist	

Catalog No.	Product Name	Primary Function	
4019	JNJ 10181457	H ₃ antagonist	
4020	JNJ 5207852	High affinity H ₃ antagonist	
2034	ROS 234	Potent H ₃ antagonist	
4441	SEN 12333	${ m H_3}$ antagonist; also ${ m lpha7}$ nAChR agonist	
2493	<u>VUF 5681</u>	Potent H ₃ receptor silent antagonist	
H ₄ Receptors			
Agonists			
2342	4-Methylhistamine	Selective, high affinity H ₄ agonist	
4769	<u>VUF 10460</u>	Selective H ₄ agonist	
2494	<u>VUF 8430</u>	Potent, high affinity H ₄ agonist	
Antagonists			
3753	<u>A 943931</u>	Potent and selective H ₄ antagonist	
3640	<u>A 987306</u>	Potent and selective H ₄ antagonist	
2441	JNJ 10191584	Selective H ₄ antagonist; orally active	
6279	JNJ 39758979	High affinity and selective H ₄ antagonist; orally bioavailable	
4021	JNJ 7777120	Selective H ₄ antagonist	
Non-selective Histamine Compounds			
Agonists			
3545	<u>Histamine</u>	Endogenous histamine agonist	
0646	HTMT	H ₁ and H ₂ agonist	
0729	<u>Imetit</u>	Standard H_3 and H_4 agonist $(H_3 > H_4)$	
0932	<u>Immepip</u>	$\rm H_3$ and $\rm H_4$ agonist	
Antagonists			
0752	Clobenpropit	Highly potent H ₃ antagonist; also H ₄ partial agonist	
0644	Thioperamide	H ₃ antagonist; also H ₄ inverse agonist	
Histaminergic-related Compounds			
4857	Amlexanox	Inhibits histamine release from rat mast cells; anti-allergic agent	
3201	Dimebon	Non-selective antihistamine; displays neuroprotectant and cognitive enhancing abilities	
0743	DPPE	Inhibitor of histamine binding at the intracellular binding site	
3848	Sinomenine	Anti-inflammatory; causes mast cell degranulation and histamine release	
0512	SKF 91488	Histamine N-methyltransferase inhibitor	
6424	Sodium Cromoglicate	Inhibits histamine release and mast cell degranulation	

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- Neurodegeneration
- G Protein-Coupled Receptors
- Pain Research
- Ion Channels



Life Science Posters

- Multiple Sclerosis
- Pain
- · Parkinson's Disease
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