IMIDAZOLE DERIVATIVES AND THEIR USE AS CYTOKINE INHIBITORS

Novel 2,4,5-arylimidazole compounds and compositions for use in therapy.
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IMIDAZOLE DERIVATIVES AND THEIR USE AS CYTOKINE INHIBITORS

This invention relates to a novel group of imidazole compounds, processes for the preparation thereof, the use thereof in treating cytokine mediated diseases and pharmaceutical compositions for use in such therapy.

BACKGROUND OF THE INVENTION:

There has been much interest in the past few years in compounds which are cytokine-inhibitors, for use in treating disease states which are associated with the excessive or unregulated production of cytokines. Compounds of the general formula (A):

\[
\begin{array}{c}
\text{R}_a \\
N \equiv W \\
\text{R}_b \\
\end{array}
\]

(A)

wherein \( \text{R}_a \) is pyridyl, \( \text{R}_b \) is optionally substituted phenyl and \( W \) is a partially or fully unsaturated fused 5- or 6-membered heterocyclic ring, such as pyrrolyl, pyridyl, dihydropyrrrol, dihydropyridinyl, dihydrothiazolyl or tetrahydro-triazinyl, are inhibitors of the cytokines IL-1 and TNF (see WO88/01169, WO90/15534, WO91/00992, WO92/10190, WO92/10498 and WO92/12154, published after the filing of this application). In addition, these compounds are also inhibitors of the enzyme 5-lipoxygenase. We have now surprisingly found that if the ring \( W \) is replaced by certain substituents at the 2-position, cytokine-inhibitory activity is maintained. Such compounds are generically 2-substituted-4-aryl-5-heteroaryl-imidazoles. Compounds within this class have already been extensively investigated, as anti-inflammatory agents, acting principally as cyclo-oxygenase inhibitors, as described in, for instance, US patents 3,707,405 and 3,929,807. The latter discloses compounds of the general formula (B):

\[
\begin{array}{c}
\text{R}_a \\
N \equiv \text{R}_c \\
\text{R}_b \\
\end{array}
\]

(B)

wherein one of \( \text{R}_a \) and \( \text{R}_b \) is optionally substituted phenyl and the other is a 6-membered heterocyclic ring with 1 or 2 nitrogen atoms and \( \text{R}_c \) represents lower alkyl, cycloalkyl or phenyl optionally substituted by halogen, lower alkyl or lower alkoxy, in particular the compound 2-(4-chlorophenyl)-4-(4-methoxyphenyl)-5-(4-pyridyl)-imidazole. These compounds are said to have
anti-inflammatory, analgesic and antipyretic activity. There is however no mention that these compounds may be cytokine inhibitors.

FULL DESCRIPTION OF THE INVENTION:

Accordingly, the present invention provides a compound of formula (I):

![Chemical Structure](image)

wherein:

- **R**₁ is 4-pyridyl, pyrimidinyl, quinolyl, isoquinolinyl, 1-imidazolyl or 1-benzimidazolyl which is optionally substituted with one or two substituents each of which is independently selected from C₁-4 alkyl, halo, C₁-4 alkoxy, C₁-4 alkylthio, NH₂, mono- or di-C₁-6-alkylamino or N-heterocyclyl ring which ring has from 5 to 7 members and optionally contains an additional heteroatom selected from oxygen, sulfur or NR₂;

- **R**₂ is R₈ or -OR₁₂;

- **R**₃ is -XₐP(Z)(XₐR₁₃)₂ or Q-(Y₁)ₜ ;

- **Q** is an aryl or heteroaryl group;

- **t** is an integer of 1 to 3;

- **Xₐ** is -NR₆-, -O-, -S- or a C₁-10 alkylen chain optionally substituted by C₁-4 alkyl and optionally interrupted by -NR₆-, -O- or -S-;

- **Xₐ** is -CR₁₀R₂₀ₙ-, -NR₆-, -O- or -S-;

- **Z** is oxygen or sulfur;

- **n** is 0 or an integer from 1 to 10;

- **Y₁** is independently selected from hydrogen, C₁-5 alkyl, halo-substituted C₁-5 alkyl, halon, -XₐP(Z)-(XₐR₁₃)₂ or -(CR₁₀R₂₀ₙ)Y₂;

- **Y₂** is -OR₈₆, -NO₂, -S(O)mR₁₁, -SR₈₆, -S(O)mOR₈₆, -S(O)mNR₆R₉₆, -NR₆R₉₆, -O(CR₁₀R₂₀ₙ)ₙNR₆Rₙ₉₆, -C(O)R₈₆, -CO₂R₈₆, -CO₂(CR₁₀R₂₀ₙ)m'-CONR₆R₉₆, -Z(C)R₈₆, -CN, -C(Z)NR₆R₉₆, -NR₁₀(C(Z))R₉₆, -C(Z)NR₆R₉₆, -NR₁₀(C(Z))NR₆R₉₆, -NR₁₀O(C(Z))R₉₆, -N(O(R₂₁)C(Z))NR₆R₉₆, -N(O(R₂₁)C(Z))R₉₆, -C(=NOR₂₁)R₈₆, -NR₁₀C(=NR₁₅)SR₁₁, -NR₁₀C(=NR₁₅)NR₆R₉₆, -NR₁₀C(=CR₁₄R₂₄)SR₁₁, -NR₁₀C(=CR₁₄R₂₄)NR₆R₉₆, -NR₁₀C(O)(O)NR₆R₉₆, -NR₁₀C(O)(O)OR₁₀, -C(=NR₁₃)NR₆R₉₆, -C(=NR₁₃)NR₆R₉₆, -C(=NR₁₃)ZR₁₁, -OC(Z)NR₆R₉₆, -NR₁₀S(O)mCF₃, -NR₁₀C(Z)OR₁₀, 5-(R₁₈)-1,2,4-oxadiazolo-3-yl or 4-(R₁₂)-5-(R₁₈R₁₉)-4,5-dihydro-1,2,4-oxadiazol-3-yl;

- **m'** is 1 or 2;
n' is an integer from 1 to 10;
R_4 is phenyl, naphth-1-y1 or naphth-2-y1 which is optionally substituted by
one or two substituents, each of which is independently selected, and
which, for a 4-phenyl, 4-naphth-1-y1 or 5-naphth-2-y1 substituent, is halo,
cyano, -C(Z)NR_7R_17, -C(Z)OR_23, -(CR_10R_20)_mCOR_36, -SR_5, -SOR_5, -OR_36,
halo-substituted-C_1-4 alkyl, C_1-4 alkyl, -ZC(Z)R_36, -NR_10C(Z)R_23, or
-(CR_10R_20)_mNR_10R_20 and which, for other positions of substitution, is
halo, cyano, -(C)NR_16R_26, -(C)OR_8, -(CR_10R_20)_mCOR_8, -S(O)_mR_8, -OR_8,
halo-substituted-C_1-4 alkyl, -C_1-4 alkyl, -(CR_10R_20)_mNR_10C(Z)R_8,
-NR_10S(O)_mR_11, -NR_10S(O)_mNR_7R_17 wherein m is 1 or 2, -ZC(Z)R_8 or
-(CR_10R_20)_mNR_16R_26;

m is 0, or the integer 1 or 2;
R_5 is hydrogen, C_1-4 alkyl, C_2-4 alkenyl, C_2-4 alkylnyl or NR_7R_17, excluding
the moieties -SR_5 being -SNR_7R_17 and -SOR_5 being -SOH;

R_6 is C_1-4 alkyl, halo-substituted-C_1-4 alkyl, C_2-4 alkenyl, C_2-4 alkylnyl or C_3-5
cycloalkyl;

R_7 and R_17 is each independently selected from hydrogen or C_1-4 alkyl or R_7
and R_17 together with the nitrogen to which they are attached form a
heterocyclic ring of 5 to 7 members which ring optionally contains an
additional heteroatom selected from oxygen, sulfur or NR_22;

R_8 is hydrogen, heterocyclyl, heterocyclylalkyl or R_11;

R_9 is hydrogen, C_1-10 alkyl, C_2-10 alkenyl, C_2-10 alkylnyl, C_3-7 cycloalkyl,
C_5-7 cycloalkenyl, aryl, arylalkyl, heteroaryl or heteroarylalkyl or R_8 and
R_9 may together with the nitrogen to which they are attached form a
heterocyclic ring of 5 to 7 members which ring optionally contains an
additional heteroatom selected from oxygen, sulfur or NR_12;

R_10 and R_20 is each independently selected from hydrogen or C_1-4 alkyl;
R_11 is C_1-10 alkyl, halo-substituted C_1-10 alkyl, C_2-10 alkenyl, C_2-10
alkynyl, C_3-7 cycloalkyl, C_5-7 cycloalkenyl, aryl, arylalkyl, heteroaryl or
heteroarylalkyl;

R_12 is hydrogen, -(C)R_13 or optionally substituted C_1-4 alkyl, optionally
substituted aryl or optionally substituted aryl-C_1-4 alkyl;

R_13 is hydrogen, C_1-10 alkyl, cycloalkyl, heterocyclyl, aryl, arylalkyl,
heteroaryl or heteroarylalkyl;

R_14 and R_24 is each independently selected from hydrogen, alkyl, nitro or
cyano; R_15 is hydrogen, cyano, C_1-4 alkyl, C_3-7 cycloalkyl or aryl;
R_16 and R_26 is each independently selected from hydrogen or optionally
substituted C_1-4 alkyl, optionally substituted aryl or optionally
substituted aryl-C_1-4 alkyl, or together with the nitrogen which they are
attached form a heterocyclic ring of 5 to 7 members which ring optionally contains an additional heteroatom selected from oxygen, sulfur or NR\\(_2\); R\(_{18}\) and R\(_{19}\) is each independently selected from hydrogen, C\(_{1-4}\) alkyl, substituted alkyl, optionally substituted aryl, optionally substituted arylalkyl or together denote a oxygen or sulfur;

R\(_{21}\) is hydrogen, a pharmaceutically acceptable cation, C\(_{1-10}\) alkyl, C\(_3\)-7 cycloalkyl, aryl, aryl C\(_{1-4}\) alkyl, heteroaryl, heteroarylalkyl, heterocyclyl, aroyl, or C\(_{1-10}\) alkoyl; R\(_{22}\) is R\(_{10}\) or C(Z)-C\(_{1-4}\) alkyl;

R\(_{23}\) is C\(_{1-4}\) alkyl, halo-substituted-C\(_{1-4}\) alkyl, or C\(_3\)-5 cycloalkyl; R\(_{96}\) is hydrogen or R\(_{23}\); or a pharmaceutically acceptable salt thereof; and excluding 2-(4-chlorophenyl)-4-(4-methoxyphenyl)-5-(4-pyridyl)imidazole.

Suitable R\(_1\) moieties include 4-pyridyl, 4-pyrimidinyl, 4-quinolyl, 6-isoquinolinyl, 1-imidazolyl and 1-benzimidazolyl, of which 4-pyridyl, 4-pyrimidinyl and 4-quinolyl, especially 4-pyridyl, are preferred. A preferred substituent for all R\(_1\) moieties is C\(_{1-4}\) alkyl, in particular methyl. More preferred as a substituted R\(_1\) moiety is the 4-pyridyl derivative substituted at the 2-position with C\(_{1-4}\) alkyl, especially 2-methyl-4-pyridyl. Also preferred is the 4-pyrimidinyl derivative substituted at the 2-position with C\(_{1-4}\) alkyl or NR\(_{10}\)R\(_{20}\).

Preferably, R\(_2\) is hydrogen or C\(_{1-10}\) alkyl, more preferably, hydrogen or methyl.

Preferably, the R\(_3\) moieties is an (un)substituted aryl or heteroaryl moiety Q, also referred to as Q-(Y\(_1\))\(_t\). Preferably, when Q is an aryl, specifically phenyl, and when Q is a heteroaryl, preferred groups include pyrrole, pyridine, or pyrimidine. More preferred is Q as phenyl. All preferred moieties are independently substituted by (Y\(_1\))\(_t\), wherein t is an integer of 1 to 3. Preferably t is 1 or 2. More preferably, when R\(_3\) is monosubstituted phenyl, the substituent is located at the 4-position.

Suitably the aryl or heteroaryl moiety of Q is substituted by up to three substituents Y\(_1\) each of which is independently selected from C\(_{1-5}\) alkyl, halo-substituted C\(_{1-5}\) alkyl, halogen, -X\(_a\)-P(Z)-(X\(_b\)R\(_{13}\))\(_2\) or -(CR\(_{10}\)R\(_{20}\))\(_n\)Y\(_2\) wherein Y\(_2\) is -OR\(_8\), -NO\(_2\), -S(O)mR\(_{11}\), -SR\(_8\), -S(O)m'OR\(_8\), -S(O)mNR\(_8\)R\(_9\), -NR\(_8\)R\(_9\), -O(CR\(_{10}\)R\(_{20}\))\(_n\)NR\(_8\)R\(_9\), -C(O)R\(_8\), -CO\(_2\)R\(_8\), -C(O)R\(_8\), -CO\(_2\)R\(_8\), -CONR\(_8\)R\(_9\), -ZC(O)R\(_8\), -CN, -C(Z)NR\(_8\)R\(_9\), -NR\(_{10}\)C(Z)R\(_8\), -C(Z)NR\(_8\)OR\(_9\), -NR\(_{10}\)C(Z)NR\(_8\)R\(_9\), -NR\(_{10}\)S(O)mR\(_{11}\), -N(OH\(_2\))C(Z)NR\(_8\)R\(_9\), -N(OH\(_2\))C(Z)R\(_8\), -C(=NOR\(_2\))R\(_8\),
-NR_{10}C(=NR_{15})SR_{11}, -NR_{10}C(=NR_{15})NR_{8}R_{9}, -NR_{10}C(=CR_{14}R_{24})SR_{11},
-NR_{10}C(=CR_{14}R_{24})NR_{8}R_{9}, -NR_{10}C(O)(O)NR_{8}R_{9}, -NR_{10}C(O)(O)OR_{10},
-C(=NR_{13})NR_{8}R_{9}, -C(=NOR_{13})NR_{8}R_{9}, -C(=NR_{13})ZR_{11}, -OC(Z)NR_{8}R_{9},
-NR_{10}S(O)_{m}CF_{3}, -NR_{10}C(Z)OR_{10}, 5-(R_{18})-1,2,4-oxadiazol-3-yl or 4-(R_{12})-5-
(R_{18}R_{19})-4,5-dihydro-1,2,4-oxadiazol-3-yl; m' is 1 or 2; R_{9} is hydrogen, C_{1-10}
alkyl, C_{2-10} alkenyl, C_{2-10} alkynyl, C_{3-7} cycloalkyl, C_{5-7} cycloalkenyl, aryl,
arlylalkyl, heteroaryl or heteroaryllalkyl or R_{8} and R_{9} may together with the
nitrogen to which they are attached form a heterocyclic ring of 5 to 7
members which ring optionally contains an additional heteroatom selected
from oxygen, sulfur or NR_{12}; R_{14} and R_{24} is each independently selected from
hydrogen, alkyl, nitro or cyano; R_{15} is hydrogen, cyano, C_{1-4} alkyl, C_{3-7}
cycloalkyl or aryl; R_{18} and R_{19} is each independently selected from hydrogen,
C_{1-4} alkyl, substituted alkyl, optionally substituted aryl, arylalkyl or together
with the carbon to which they are attached denote a double bonded oxygen or
sulfur, i.e., a C=O or C=S; and R_{21} is hydrogen, a pharmaceutically acceptable
cation, alkyl, cycloalkyl, aryl, arylalkyl, heteroaryl, heterocyclic,
heteroaryllalkyl, aryl, alkyl.

Preferred substituents Y_{1} for use in R_{3} include halogen, C_{1-5} alkyl
and -(CR_{10}R_{20})_{n}Y_{2} wherein Y_{2} is -OR_{8}, -NO_{2}, -SO(O)_{m}R_{11}, -SR_{8},
-S(O)_{m}NR_{8}R_{9}, -NR_{8}R_{9}, -O(CR_{10}R_{20})_{n}NR_{8}R_{9}, -C(O)R_{8}, -CO_{2}R_{8},
-CO_{2}(CR_{10}R_{20})_{n}CONR_{8}R_{9}, -CN, -C(Z)NR_{8}R_{9}, -NR_{10}S(O)_{m}R_{11},
-NR_{10}C(Z)R_{8}, -NR_{10}C(Z)NR_{8}R_{9}, -C(Z)NR_{8}OR_{9}, -N(OR_{21})C(Z)NR_{8}R_{9},
-NR_{10}C(=NR_{15})NR_{8}R_{9}, -C(=NOR_{13})NR_{8}R_{9}, 5-(R_{18})-1,2,4-oxadizaol-3-yl and
4-(R_{12})-5-(R_{18}R_{19})-4,5-dihydro-1,2,4-oxadiazol-3-yl.

Preferred substituents Y_{1} for use in R_{3} when the aryl or heteroaryl
group Q is mono-substituted include -(CR_{10}R_{20})_{n}Y_{2} wherein: n is 0, 1, 2 or
3, preferably 0 or 1; and Y_{2} is -OR_{8}, especially where R_{8} is hydrogen or
C_{1-10} alkyl; -NO_{2}, -SO(O)_{m}R_{11}, especially where R_{11} is C_{1-10} alkyl; -SR_{8},
especially where R_{8} is C_{1-10} alkyl; -S(O)_{m}NR_{8}R_{9}, especially where R_{8} and
R_{9} is each hydrogen or C_{1-10} alkyl or R_{8} and R_{9} together with the nitrogen
to which they are attached form a 5 to 7 membered ring which optionally
includes another heteroatom selected from oxygen, sulfur or NR_{12} and m is
2; n' is 1 to 10; -NR_{8}R_{9}, especially where R_{8} and R_{9} is each hydrogen,
methyl or benzyl or R_{8} and R_{9} together with the nitrogen to which they are
attached form a 5 to 7 membered ring which optionally includes another
heteroatom selected from oxygen, sulfur or NR_{12}; -O(CR_{10}R_{20})_{n}NR_{8}R_{9},
especially where R_{8} and R_{9} is each C_{1-10} alkyl; -C(O)R_{8}, especially where
R_{8} is hydrogen or C_{1-10} alkyl; -CO_{2}R_{8}, especially where R_{8} is hydrogen or
C_{1-10} alkyl; -CO_{2}(CR_{10}R_{20})_{n}CONR_{8}R_{9}, especially where R_{8} and R_{9} is
hydrogen or C₁₋₁₀ alkyl; -CN; -C(Z)NR₈R₉, especially where R₈ and R₉ is hydrogen or C₁₋₁₀ alkyl; -NR₁₀S(O)ₘR₁₁, especially where R₁₀ is hydrogen or C₁₋₁₀ alkyl and R₁₁ is C₁₋₁₀ alkyl or a halosubstituted; -NR₁₀C(Z)R₈, especially where R₈ is C₁₋₁₀ alkyl and R₁₀ is hydrogen and Z is oxygen;

-C(Z)NR₈₉OR₉, especially where R₈ and R₉ is each hydrogen and Z is oxygen;
-NR₁₀C(Z)NR₈R₉, especially where R₈ and R₉ is each hydrogen or C₁₋₁₀ alkyl and Z is oxygen; -N(OH)₂C(Z)NR₈₉R₉, especially where R₈ especially where R₈, R₉ and R₂₁ is each hydrogen or C₁₋₁₀ alkyl and Z is oxygen;

-C(=NOR₁₃)NR₈R₉, especially where R₈, R₉ and R₁₃ is each hydrogen;
-NR₁₀C(=NHR₁₅)NR₈R₉, especially where R₈ and R₉ is hydrogen, C₁₋₁₀ alkyl or arylalkyl and R₁₅ is cyano; and 5-(R₁₈)₁₂,₂₄-oxadiazol-3-yl and 4-(R₁₂)₅-(R₁₈R₁₉)-4,₅-dihydro-₁,₂₄-oxadiazol-3-yl, especially where R₁₂ is hydrogen and R₁₈ and R₁₉ is each hydrogen or C₁₋₁₀ alkyl or together are oxo.

Preferred substituents for use in R₃ when the aryl or heteroaryl group Q is disubstituted include those hereinbefore listed for use when Q is monosubstituted and, as further substituent(s), halogen and C₁₋₁₀ alkyl. When R₃ is phenyl substituted with two or three substituents, the alkyl moieties preferably have from one to three carbons, more preferably one. Preferred ring positions for two substituents are the 3- and 4-positions and, for three substituents, the 3-, 4- and 5-positions. The substituent at the 3- and 5-positions is preferably C₁₋₂ alkyl, such as methyl, or halogen, such as bromo, fluoro or chloro, while the substituent at the 4-position is preferably hydroxyl.

More preferably, for R₃ substituents wherein Y₁ is (CR₁₀R₂₀)n, n is 0 or 1 and Y₂ is -OH, -S(O)ₘR₁₁, especially where R₁₁ is C₁₋₁₀ alkyl; -SR₈, especially where R₈ is C₁₋₁₀ alkyl; -NR₈R₉, especially where R₈ and R₉ is hydrogen, alkyl, aryl alkyl, or aryl or R₈ and R₉ together with the nitrogen to which they are attached form a pyrrolidinyl, piperidinyl or morpholinyl ring, more preferably the R₈ and R₉ terms in the NR₈R₉ moiety are hydrogen, methyl or benzyl; -CO₂R₈, especially where R₈ is hydrogen or C₁₋₁₀ alkyl; -S(O)ₘNR₈R₉, especially where R₈ and R₉ is each hydrogen or C₁₋₁₀ alkyl; -NR₁₀S(O)ₘR₁₁, especially where R₁₀ is hydrogen and R₁₁ is C₁₋₁₀ alkyl or 5-(R₁₈)₁₂,₂₄-oxadiazol-3-yl and 4-(R₁₂)₅-(R₁₈R₁₉)-4,₅-dihydro-₁,₂₄-oxadiazol-3-yl, especially where R₁₂ is hydrogen and R₁₈ and R₁₉ is hydrogen or C₁₋₁₀ alkyl or together are oxo.

Most preferably, Y₁ is methylthio, ethylthio, methylsulfanyl, ethylsulfanyl, methylsulfonyl, N,N-dimethylaminomethyl, N-benzyl-N-
methylaminomethyl, N-morpholinomethyl, methanesulfonamido, sulphonamidomethyl, 5-methyl-4,5-dihydro-1,2,4-oxadiazol-3-yl or 5,5-dimethyl-4,5-dihydro-1,2,4-oxadiazol-3-yl.

In all instances herein where there is an alkenyl or alkynyl moiety as a substituent group, such as in R₅, R₈, R₉, or R₁₁ the unsaturated linkage, i.e., the vinylene or acetylene linkage is preferably not directly attached to the nitrogen, oxygen or sulfur moieties, for instance in Y₂ as C(Z)NR₈OR₉, NR₁₀C(Z)NR₈R₉, or OR₉. As used herein, "optionally substituted" unless specified refers to such groups as halogen, hydroxyl, alkoxy, S(O)m alkyl, amino, mono & di-substituted amino, such as a NR₇R₁₇ group, alkyl or cycloalkyl, i.e. such as in optionally substituted aryl or optionally substituted arylalkyl.

When R₃ includes a X₄-P(Z)(X₅R₁₃)₂ group linked either directly to the imidazole ring or indirectly via an aryl or heteroaryl group, X₄ is suitably oxygen or C₁₋₄ alkylene, optionally interrupted by oxygen, for instance -CH₂OCH₂- and Z and X₅ is each oxygen, such that the preferred groups include -OP(O)(OR₁₃)₂ and -CH₂OCH₂-P(O)(OR₁₃)₂.

Preferred substitutions for R₄ when this is a 4-phenyl, 4-naphth-1-yl or 5-naphth-2-yl moiety are one or two substituents each independently selected from halogen, -SR₅, -SOR₅, -OR₃₆, or -(CR₁₀R₂₀)mNR₁₀R₂₀, and for other positions of substitution on these rings preferred substitution is halogen, -S(O)mR₈, -OR₈, -(CR₁₀R₂₀)mNR₁₆R₂₆, -NR₁₀C(Z)R₈ and -NR₁₀S(O)mR₁₁. More preferred substituents for the 4-position in phenyl and naphth-1-yl and on the 5-position in naphth-2-yl include halogen, especially fluoro and chloro, and -SR₅ and -SOR₅ wherein R₅ is preferably a C₁₋₂ alkyl, more preferably methyl; of which fluoro is especially preferred. Preferred substituents for the 3-position in phenyl and naphth-1-yl include: halogen, especially chloro; -OR₈, especially C₁₋₄ alkoxy; amino; -NR₁₀C(Z)R₈, especially -NHCO(C₁₋₁₀ alkyl); and -NR₁₀S(O)mR₁₁, especially -NHSO₃(C₁₋₁₀ alkyl). Preferably, the R₄ moiety is an unsubstituted or substituted phenyl moiety. More preferably, R₄ is phenyl or phenyl substituted at the 4-position with fluoro and/or substituted at the 3-position. In a preferred subgenus of compounds of formula (I), R₁ is 4-pyridyl, 2-alkyl-4-pyridyl or 4-quinolyl; R₂ is hydrogen or methyl; R₃ is phenyl or phenyl substituted, preferably at the 4-position, with a substituent selected from -(CR₁₀R₂₀)nY₂ wherein Y₂ is wherein n is 0, 1, 2 or 3 and Y₂ is -OR₈, -NO₂, -S(O)mR₁₁, -SR₈, -S(O)mNR₈R₉, -NR₈R₉, -O(CR₁₀R₂₀)mNR₈R₉,
-C(O)R₈, -CO₂R₈, -CO₂(CR₁₀R₂₀)ₙCONR₆R₉, -CN, -C(Z)NR₆R₉,
-C(Z)NR₈OR₉, -NR₁₀S(O)ₙR₁₁, -NR₁₀C(Z)R₈, -NR₁₀C(Z)NR₆R₉,
-C(≡NOR₁₃)NR₆R₉, -NR₁₀C(≡CR₁₄R₂₄)NR₆R₉, 5-(R₁₈)-1,2,4-oxadiazol-3-yl,
4-(R₁₂)-5-(R₁₈R₁₉)-4,5-dihydro-1,2,4-oxadiazol-3-yl, a 3,5-dimethyl or
dibromo-4-hydroxy grouping; and R₄ is phenyl or phenyl substituted at the
4-position with fluoro and/or substituted at the 3-position with fluoro, chloro,
C₁₄ alkoxy, methanesulfonamido or acetamido.

In a more preferred subgenus, R₁ is 4-pyridyl, 2-methyl-4-pyridyl or
4-quinolyl; R₂ is hydrogen or methyl; R₃ is phenyl substituted at the 4-
position with C₁₀ alklythio, C₁₀ alklysulfanyl, C₁₀ alkylsulfonyl,
N,N-di(C₁₀ alkyl)amino C₁₂ alkyl, N-aralkyl-N-C₁₀ alkylamino C₁₂
alkyl, N-morpholino C₁₂ alkyl, C₁₀ alkylsulfonamido, sulphonamido C₁₂
alkyl, 5-C₁₀ alkyl-4,5-dihydro-1,2,4-oxadiazol-3-yl or 5,5-di(C₁₀ alkyl)-
4,5-dihydro-1,2,4-oxadiazol-3-yl; and R₄ is phenyl or phenyl substituted at
the 4-position with fluoro and/or substituted at the 3-position with fluoro,
chloro, C₁₄ alkoxy, methane-sulfonamido or acetamido.

Suitable pharmaceutically acceptable salts are well known to those
skilled in the art and include basic salts of inorganic and organic acids, such
as hydrochloric acid, hydrobromic acid, sulphuric acid, phosphoric acid,
methane sulphonic acid, ethane sulphonic acid, acetic acid, malic acid,
tartaric acid, citric acid, lactic acid, oxalic acid, succinic acid, fumaric acid,
maleic acid, benzoic acid, salicylic acid, phenylactic acid and mandelic acid.
In addition, pharmaceutically acceptable salts of compounds of formula (I)
may also be formed with a pharmaceutically acceptable cation, for instance,
if a substituent Y₁ in R₃ comprises a carboxy group. Suitable pharma-
aceutically acceptable cations are well known to those skilled in the art and
include alkaline, alkaline earth, ammonium and quarternary ammonium
cations.

The following terms, as used herein, refer to:

- "halo" - all halogens, that is chloro, fluoro, bromo and iodo;
- "C₁₁₀alkyl" or "alkyl" - both straight and branched chain radicals
  of 1 to 10 carbon atoms, unless the chain length is otherwise limited,
  including, but not limited to, methyl, ethyl, n-propyl, iso-propyl, n-butyl, sec-
  butyl, iso-butyl, tert-butyl, and the like;
- "aryl" - phenyl and naphthyl;
- "heteroaryl" (on its own or in any combination, such as
  "heteroaryloxy") - a 5-10 membered aromatic ring system in which one or
  more rings contain one or more heteroatoms selected from the group
consisting of N, O or S, such as, but not limited, to pyrrole, quinoline, isoquinoline, pyridine, pyrimidine, oxazole, thiazole, thiadiazole, triazole, imidazole, or benzimidazole;
  • "heterocyclic" (on its own or in any combination, such as "heterocyclalkyl") - a saturated or wholly or partially unsaturated 4-10 membered ring system in which one or more rings contain one or more heteroatoms selected from the group consisting of N, O, or S; such as, but not limited to, pyrrolidine, piperidine, piperazine, morpholine, imidazolidine or pyrazolidine;
  • "aryl" - a C(=O)Ar, wherein Ar is as phenyl, naphthyl, or aryl alkyl derivatives, such as benzyl and the like;
  • "alkoyl" - a C(O)C_{1-10}alkyl wherein the alkyl is as defined above;
  • "sulfinyl" - the oxide (SO) of the corresponding sulfide whilst the term "thio" refers to the sulfide.

The compounds of the present invention may contain one or more asymmetric carbon atoms and may exist in racemic and optically active forms. All of these compounds are included within the scope of the present invention.

For the purposes herein of nomenclature, the compounds of formula (I) are named by their position corresponding to:

![Chemical structure](image)

Especially preferred compounds of formula (I) include:

4-(4-Fluorophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole;
4-(4-Fluorophenyl)-2-(4-ethyliophenyl)-5-(4-pyridyl)imidazole;
4-(4-Fluorophenyl)-2-(4-methylsulfonylphenyl)-5-(4-pyridyl)imidazole;
4-(4-Fluorophenyl)-2-(4-methylsulfinylphenyl)-5-(4-pyridyl)imidazole;
4-(4-Fluorophenyl)-2-(4-ethylsulfinylphenyl)-5-(4-pyridyl)imidazole;
4-(3-Chlorophenyl)-2-(4-methylsulfinylphenyl)-5-(4-pyridyl)imidazole;
2-[4-(N-Methyl-N-benzyl)aminomethylphenyl]-4-(4-fluorophenyl)-5-(4-pyridyl)imidazole;
4-(4-Fluorophenyl)-5-[4-(2-methylpyridyl)]-2-(4-methylthiophenyl)imidazole;
4-(4-Fluorophenyl)-5-[4-(2-methylpyridyl)]-2-(4-methylsulfinylphenyl)imidazole;
4-(4-Fluorophenyl)-2-(4-methylsulfinylphenyl)-5-(4-quinolyl)imidazole;
2-[4-(N-Morpholino)methylphenyl]-4-(4-fluorophenyl)-5-(4-pyridyl)-imidazole; and
pharmacologically acceptable salts thereof.

Other preferred compounds of formula (I) include:

2-[(4-N,N-Dimethyl)aminomethylphenyl]-4-(4-fluorophenyl)-5-(4-pyridyl)-imidazole;

2-(4-Methanesulfonamidophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)imidazole;

4-(4-Fluorophenyl)-2-(4-methylthiophenyl)-5-(4-quinolyl)imidazole;

4-(3-Chlorophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole;

4-(3-Methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole;

4-(3-Methoxyphenyl)-2-(4-methylsulfonylphenyl)-5-(4-pyridyl)imidazole;

4-(3-Methanesulfonamidophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole;

4-(3-Methanesulfonamidophenyl)-2-(4-methylsulfonylphenyl)-5-(4-pyridyl)imidazole;

3-[4-(4-Fluorophenyl)-5-(4-pyridyl)imidazol-2-yl]phenyl-5,5-dimethyl-4,5-
dihydro-1,2,4-oxadiazole; or

3-[4-(4-Fluorophenyl)-5-(4-pyridyl)imidazol-2-yl]phenyl-5-methyl-4,5-
dihydro-1,2,4-oxadiazole; and
pharmacologically acceptable salts thereof.

Compounds of formula (I) are imidazole derivatives which may be readily prepared using procedures well-known to those skilled in the art, and described in, for instance, Comprehensive Heterocyclic Chemistry, ed Katritzky and Rees, Pergamon Press, 1984, 5, 457-497, from starting materials which are either commercially available or can be prepared from such by analogy with well-known processes. A key step in many such syntheses is the formation of the central imidazole nucleus, to give compounds of formula (I). Suitable procedures are described in inter alia US patent nos. 3,707,475 and 3,940,486 which are herein incorporated by reference in their entirety. These patents describe the synthesis of α-diketones and α-hydroxyketones (benzoin) and their subsequent use in preparing imidazoles and N-hydroxyl imidazoles. Thereafter, further compounds of formula (I) may be obtained by manipulating substituents in any of the groups R₁, R₂, R₃ and R₄ using conventional functional group interconversion procedures.

In particular, in a first process, compounds of formula (I) may be prepared by condensing an α-diketone of formula (II):
R₁COCOR₄

wherein R₁ and R₄ are as hereinbefore defined, or an equivalent thereof, with an aldehyde of the formula (III):

R₃CHO

wherein R₃ is as hereinbefore defined, or an equivalent thereof, and, if necessary, with ammonia or a source thereof, under imidazole-ring forming conditions.

Suitable equivalents of the α-diketone are well known to those skilled in the art and include the corresponding α-keto-oxime and α-dioxime.

Suitable equivalents of the aldehyde of formula (III) are well known in the art and include the corresponding oxime and acetal.

Ammonia, or a source thereof, is preferably used in excess, with at least a dimolar amount being used in the case of the α-diketone and at least an equimolar amount in the case of the α-keto-oxime.

Suitable sources of ammonia include ammonium salts of organic carboxylic acids, such as an ammonium C₁₋₆ alkanolate, for instance ammonium acetate and ammonium formate, preferably ammonium acetate, and carboxylic amides, in particular of formic acid, such as formamide. An ammonium salt is generally used in large excess and in the presence of an acid, such as a C₁₋₆ carboxylic acid which acid may also be used as a solvent for the reaction. If formamide is used, this may be used in excess, as the reaction solvent. An alternative solvent such as ethanol or dimethyl sulfoxide (Lantos et al, J Het Chem, 19, 1375, 1982) may be used. An additional solvent may also be employed, for instance, dimethyl formamide may be used with formamide. The reaction is generally carried out at elevated temperatures, for instance under reflux conditions, and if desired, in a sealed vessel optionally under pressure and/or an inert gas atmosphere, for instance nitrogen.

A further suitable source of ammonia is hydroxylamine, in which case the initially formed imidazole is an N-hydroxy-N-oxide imidazole. This may then be reduced to the corresponding N-hydroxy imidazole by treating with a suitable reducing agent such as sodium borohydride, in an appropriate solvent such as methanol, following the method of Akange and Allan, Chem and Ind, 5 Jan 1975, 38. The N-hydroxy imidazole may in turn be converted to an imidazole of formula (I) in which R₂ is hydrogen by treatment with a conventional deoxygenating agent such as phosphorus trichloride or a trialkylphosphite such as trimethyl- or triethyl-phosphite. N-hydroxy-N-oxide imidazoles may be readily obtained by treating an α-diketone of
formula (II) with an aldehyde of formula (II) with about two equivalents of hydroxylamine or the corresponding aldoxime and about one equivalent of hydroxylamine, under proton catalysis. Alternatively, the N-oxide may be obtained by the acid catalysed condensation of the corresponding α-dioxime or α-keto-oxime with an aldoxime of the aldehyde of formula (III).

When the compound of formula (II) is an α-keto-oxime derivative, it will be appreciated that the product initially obtained will be a compound of formula (I) in which \( R_2 \) is hydroxyl which may be converted into a compound of formula (I) in which \( R_2 \) is hydrogen as described above.

It will be appreciated by those skilled in the art that in some instances, it will not be necessary to provide a separate source of ammonia as the α-diketone or aldehyde equivalent may already contain such a source. Examples of this include α-dioxime or α-keto-oxime and aldoxime.

The compounds of formula (II) may be obtained by applying well-known synthetic procedures, some of which are illustrated in schemes I and II. Although these illustrate syntheses in which \( R_4 \) is either 4-pyridyl or 4-quinolyl, they may be equally applied to any of the other heteroaryl rings within the definition of \( R_4 \) by appropriate choice of starting material.

In Scheme I, the anion prepared from 1, by treatment with a strong base such as lithium di-iso-propylamide, is condensed with a substituted benz-aldehyde, to give, after removal of the protecting group, the diol 2. This may then be converted to the α-diketone 3 by a Swern oxidation of which any number of potentially useful variations are known and may be used. The α-diketone 3 is then cyclised to an imidazole 4, a compound of formula (I), by heating 3 with a substituted benzaldehyde in a mixture of ammonium acetate, as the source of ammonia, and an appropriate solvent, for example acetic acid or DMSO. The imidazole 4 may then be transformed into other imidazoles 5 by appropriate functional group interconversion procedures. Scheme I also illustrates the preparation of a protected α-hydroxyketone 2a, by condensing the anion of 1 with an appropriately activated carbonyl derivative of a substituted benzaamide, such as the N-methoxy-N-methylamide, to yield a protected α-hydroxyketone. This adduct 2a may then be directly converted to the imidazole 5, using a combination of a copper (II) salt, such as copper (II) acetate, as an oxidising agent and ammonium acetate as a source of ammonia. The α-hydroxyketone 2a may also be deprotected and then oxidised to give an α-diketone 3, for instance using Swern oxidation.
Scheme I

Scheme II illustrates the use of an $\alpha$-keto-oxime for preparing a compound of formula (I). A heterocyclic ketone 7 is prepared by adding the anion of 4-methyl-quinoline (prepared by treatment thereof with an alkyl lithium, such as $n$-butyl lithium) to an N-alkyl-O-alkoxybenzamide. Alternatively, the anion may be condensed with a benzaldehyde, to give an alcohol which is then oxidised to the ketone 7. The $\alpha$-keto-oxime 8 is then prepared from 7 using standard conditions, such as reaction with sodium nitrite, and this may then be reacted with a benzaldehyde to afford an N-hydroxyimidazole 9, a compound of formula (I) in which $R_2$ is hydroxy. This may converted to 10, a further compound of formula (I) in which $R_2$ is hydrogen, by treating it with a deoxygenating agent such as phosphorus trichloride or a trialkyl phosphite, such as trimethyl or triethylphosphite. For compounds of formula (I) wherein $R_3$ is -(CR$_{10}$R$_{20}$)$_n$-P(Z)-(X$_b$R$_{13}$)$_2$, the reagent OHC-(CR$_{10}$R$_{20}$)$_n$-P(Z)-(X$_b$R$_{13}$)$_2$ may be used instead of OHC-C$_6$H$_4$-X to make the appropriately substituted compound 9.
Scheme II

In a further process, a compound of formula (I) may be obtained by treating an α-hydroxyketone compound of formula (IIA):

$$R'CHOHCOR''$$  \hspace{1cm} (IIA)

wherein one of $R'$ and $R''$ is $R_1$ and the other is $R_4$, a suitably protected derivative thereof or the α-hydroxy-oxime or α-haloketone derivative thereof, with an oxidising agent capable of converting said compound into the corresponding α-diketone, in the presence of an aldehyde of formula (III) or an equivalent thereof, and a source of ammonia. Suitable oxidising agents include, for example, an oxidising heavy metal salt, preferably an organic copper (II) salt, such as copper (II) acetate or copper (II) citrate. The reaction may be effected in a solvent such as acetic acid, under reflux conditions. Alternatively, a lower alkanol solvent, such as methanol or ethanol, may be used, preferably at a temperature in the region of from 30 to 100°C (see The Chemistry of Heterocyclic Compounds, Imidazole and its
derivatives, part I, ed. Weissberger, Interscience Publishers, Inc., New York, 1953, 38). This approach is also illustrated in Scheme I.

In a further process, a compound of formula (I) may be obtained by treating an amidine of formula (IV):

\[
R_3C(=\text{NH})NHR_2 \quad (IV)
\]

wherein \(R_2\) and \(R_3\) are as hereinbefore defined, or a salt thereof, with a reactive ester of an \(\alpha\)-hydroxyketone of formula (IIA) or the corresponding \(\alpha\)-haloketone, in an inert solvent such as a halogenated hydrocarbon solvent, for example chloroform, at a moderately elevated temperature and, if necessary, in the presence of a suitable condensation agent such as a base. Suitable reactive esters include esters of strong organic acids such as a lower alkane sulphonic or aryl sulphonic acid, for instance, methane or p-toluene sulphonic acid. The amidine of formula (IV) is preferably used as the salt, suitably the hydrochloride salt, which may then be converted into the free amidine in situ, by employing a two phase system in which the reactive ester is in an inert organic solvent such as chloroform, and the salt is in an aqueous phase to which a solution of an aqueous base is slowly added, in dimolar amount, with vigorous stirring. Suitable amidines of formula (IV) may be obtained by standard methods, see for instance, Garigipati R, Tetrahedron Letters, 190, 31, 1989.

In a further process, a compound of formula (I) may be obtained by treating an iminoether of formula (V):

\[
R_3C=\text{NOR} \quad (V)
\]

wherein \(R_3\) is as hereinbefore defined and \(R\) is \(C_{1-10}\) alkyl, aryl or aryl \(C_{1-4}\) alkyl, with an \(\alpha\)-amino ketone of the formula (VI):

\[
R'\text{CHNH}_2\text{COR}'' \quad (VI)
\]

wherein one of \(R'\) and \(R''\) is \(R_1\) and the other is \(R_4\) in a suitable solvent.

In a further process, \(N\)-substituted compounds of formula (I) may be prepared by treating the anion of an amide of formula (VII):

\[
R_1\text{CH}_2\text{NR}_2\text{COR}_3 \quad (VII)
\]

wherein \(R_1\) and \(R_3\) are as hereinbefore defined and \(R_2\) is as hereinbefore defined other than hydrogen, with:

(a) a nitrile of the formula (VIII):

\[
R_4\text{CN} \quad (VIII)
\]

wherein \(R_4\) is as hereinbefore defined, or

(b) an excess of an acyl halide, for instance an acyl chloride, of the formula (IX):

\[
R_4\text{COHal} \quad (IX)
\]
wherein R₄ is as hereinbefore defined and Hal is halogen, or a corresponding anhydride, to give a bis-acylated intermediate which is then treated with a source of ammonia, such as ammonium acetate.

This approach permits the regiospecific preparation of compound of formula (I) substituted at the 1-position, as illustrated in Scheme III. A primary amine RNH₂ is treated with 4-chloromethylpyridine to give 11 which is then converted to the amide 12 by standard techniques. Deprotonation of 12 with a strong amide base, such as lithium di-iso-propyl amide or sodium bis-(trimethylsilyl)amide, followed by addition of an excess of an aroyl chloride yields the bis-acylated compound 13 which is then closed to an imidazole compound of formula (I), 14, by heating in acetic acid containing ammonium acetate. Alternatively, the anion of 12 may be reacted with a substituted aryl nitrile to produce the imidazole 14 directly.

In a further process, compounds of formula (I) may be prepared by treating a compound of formula (X):

\[ R'^{'}COCHR"XₐCOR₃ \]

wherein R', R" and R₃ are as hereinbefore defined and Xₐ is O or NH, with a source of ammonia, as hereinbefore described, under imidazole ring forming conditions or cyclising the corresponding Schiff's base, formed by treating the compound of formula (X) in which Xₐ is NH with an amine R₉NH₂, for instance thermally or with the aid of a cyclising agent such as phosphorus oxychloride or phosphorus pentachloride (see Engel and Steglich, Liebig's Ann Chem, 1978, 1916 and Strzybny et al., J Org Chem, 1963, 28, 3381). Compounds of formula (X) may be obtained, for instance, by acylating the corresponding α-keto-oxime (Xₐ is NH) or α-hydroxyketone (Xₐ is O) with an
acyl halide of the formula \( R_3 COHal \) wherein \( R_3 \) is as hereinbefore defined, or the corresponding anhydride, under standard acylating conditions.

In a further process, compounds of formula (I) may be prepared by coupling a suitable derivative of a compound of formula (XI):

wherein: \( T_2 \) is a nitrogen protecting group or \( R_2 \), other than hydrogen; and \( T_1 \) is hydrogen, \( T_3 \) is \( Q \) and \( T_4 \) is \( R_4 \); \( T_1 \) is \( R_1 \), \( T_3 \) is hydrogen and \( T_4 \) is \( R_4 \); or \( T_1 \) is \( R_1 \), \( T_3 \) is \( Q \) and \( T_4 \) is hydrogen, in which \( R_1, R_2, R_3, R_4 \) and \( Q \) are as hereinbefore defined; with: (i) when \( T_1 \) is hydrogen, a suitable derivative of the heteroaryl ring \( R_1 H \), under ring coupling conditions, to effect coupling of the heteroaryl ring \( R_1 \) to the imidazole nucleus at position 5; (ii) when \( T_3 \) is hydrogen, a suitable derivative of the aryl or heteroaryl ring \( QH \), under ring coupling conditions, to effect coupling of the ring \( Q \) to the imidazole nucleus at position 2; or (iii) when \( T_4 \) is hydrogen, a suitable derivative of the aryl ring \( R_4 H \), under ring coupling conditions, to effect coupling of the aryl ring \( R_4 \) to the imidazole nucleus at position 4.

Such aryl/heteroaryl coupling reactions are well known to those skilled in the art. In general, an organometallic synthetic equivalent of an anion of one component is coupled with a reactive derivative of the second component, in the presence of a suitable catalyst. The anion equivalent may be formed from either the imidazole of formula (XI), in which case the aryl/heteroaryl compound provides the reactive derivative, or the aryl/heteroaryl compound in which case the imidazole provides the reactive derivative. Accordingly, suitable derivatives of the compound of formula (XI) or the aryl/heteroaryl rings include organometallic derivatives such as organomagnesium, organozinc, organostannane and boronic acid derivatives and suitable reactive derivatives include the the bromo, iodo, fluorosulfonate and trifluoromethanesulphonate derivatives. Suitable procedures are described in WO 91/19497, the disclosure of which is herewith incorporated.

Suitable organomagnesium and organozinc derivatives of a compound of formula (XI) may be reacted with a halogen, fluorosulfonate or triflate derivative of the heteroaryl or aryl ring, in the presence of a ring coupling catalyst, such as a palladium (O) or palladium (II) catalyst, following the procedure of Kumada et al., Tetrahedron Letters, 22, 5319 (1981). Suitable such catalysts include tetraakis-(triphenylphosphine)palladium and
PdCl₂[1,4-bis-(diphenylphosphino)-butane], optionally in the presence of lithium chloride and a base, such as triethylamine. In addition, a nickel (II) catalyst, such as Ni(II)Cl₂(1,2-biphenylphosphino)ethane, may also be used for coupling an aryl ring, following the procedure of Pridgen, J Org Chem, 1982, 47, 4319. Suitable reaction solvents include hexamethylenophosphamide. When the heteroaryl ring is 4-pyridyl, suitable derivatives include 4-bromo- and 4-iodo-pyridine and the fluorosulfonate and triflate esters of 4-hydroxy pyridine. Similarly, suitable derivatives for when the aryl ring is phenyl include the bromo, fluorosulfonate, triflate and, preferably, the iodo-derivatives. Suitable organomagnesium and organozinc derivatives may be obtained by treating a compound of formula (XI) or the bromo derivative thereof with an alkyl lithium compound to yield the corresponding lithium reagent by deprotonation or transmetallation, respectively. This lithium intermediate may then be treated with an excess of a magnesium halide or zinc halide to yield the corresponding organometallic reagent.

A trialkyltin derivative of the compound of formula (XI) may be treated with a bromide, fluorosulfonate, triflate, or, preferably, iodide derivative of an aryl or heteroaryl ring compound, in an inert solvent such as tetrahydrofuran, preferably containing 10% hexamethylenophosphamide, in the presence of a suitable coupling catalyst, such as a palladium (0) catalyst, for instance tetrakis-(triphenylphosphine)palladium, by the method described in by Stille, J Amer Chem Soc, 1987, 109, 5478, US Patents 4,719,218 and 5,002,942, or by using a palladium (II) catalyst in the presence of lithium chloride optionally with an added base such as triethylamine, in an inert solvent such as dimethyl formamide. Trialkyltin derivatives may be conveniently obtained by metllation of the corresponding compound of formula (XI) with a lithiating agent, such as s-butyl-lithium or n-butyl lithium, in an ethereal solvent, such as tetrahydrofuran, or treatment of the bromo derivative of the corresponding compound of formula (XI) with an alkyl lithium, followed, in each case, by treatment with a trialkyltin halide. Alternatively, the bromo- derivative of a compound of formula (XI) may be treated with a suitable heteroaryl or aryl trialkyl tin compound in the presence of a catalyst such as tetrakis-(triphenylphosphine)-palladium, under conditions similar to those described above.

Boronic acid derivatives are also useful. Hence, a suitable derivative of a compound of formula (XI), such as the bromo, iodo, triflate or fluorosulphonate derivative, may be reacted with a heteroaryl- or aryl-boronic acid, in the presence of a palladium catalyst such as tetrakis-(triphenylphosphine)-palladium or PdCl₂[1,4-bis-(diphenylphosphino)-

Non-aqueous conditions, for instance, a solvent such as DMF, at a temperature of about 100°C, in the presence of a Pd(II) catalyst may also be employed (see Thompson W J et al., J Org Chem, 49, 5237, 1984). Suitable boronic acid derivatives may be prepared by treating the magnesium or lithium derivative with a trialkylborate ester, such as triethyl, tri-iso-propyl or tributylborate, according to standard procedures.

In such coupling reactions, it will be readily appreciated that due regard must be exercised with respect to functional groups present in the compounds of formula (XI). Thus, in general, amino and sulfur substituents should be non-oxidised or protected and the N-1 nitrogen of a compound of formula (XI) be protected, if an NH compound is finally required. Nitro, bromo, iodo and hydroxyl groups should preferably be avoided in compounds of formula (XI) in which $T_1$ is hydrogen.

Compounds of formula (XI) are imidazoles and may be obtained by any of the procedures herein before described for preparing compounds of formula (I). In particular, an $\alpha$-halo-ketone $R_4$COCH$_2$Hal (for compounds of formula (XI) in which $T_1$ is hydrogen) or $R_1$COCH$_2$Hal (for compounds of formula (XI) in which $T_4$ is hydrogen) may be reacted with an amidine of formula (IV) or a salt thereof, in an inert solvent such as a halogenated hydrocarbon solvent, for instance chloroform, at a moderately elevated temperature, and, if necessary, in the presence of a suitable condensation agent such as a base. The preparation of suitable $\alpha$-halo-ketones is described in WO 91/19497. For a compound of formula (XI) in which $T_3$ is hydrogen, an $\alpha$-diketone of formula (II) may be condensed with a formaldehyde or an equivalent thereof, in the presence of a source of ammonia. Suitable bromo derivatives of the compound of formula (XI) may be obtained by brominating the corresponding compound of formula (XI) under standard brominating conditions, for instance bromine in a solvent such as dichloromethane or THF.

Compounds of formula (I) may also be prepared by a process which comprises reacting a compound of formula (XI), wherein $T_1$ is hydrogen, with an N-acyl heteroaryl salt, according to the method disclosed in US patents 4,803,279, 4,719,218 and 5,002,942, to give an intermediate in which the heteroaryl ring is attached to the imidazole nucleus and is present as a 1,4-dihydro derivative thereof, which intermediate may then be
subjected to oxidative-deacylation conditions. The heteroaryl salt, for instance a pyridinium salt, may be either preformed or, more preferably, prepared in situ by adding a substituted carbonyl halide (such as an acyl halide, an aryl halide, an arylalkyl haloformate ester, or, preferably, an alkyl haloformate ester, such as acetyl bromide, benzoyl chloride, benzyl chloroformate, or, preferably, ethyl chloroformate) to a solution of the compound of formula (XI) in the heteroaryl compound R1H or in an inert solvent such as methylene chloride to which the heteroaryl compound has been added. Suitable deacylating and oxidising conditions are described in U.S. Patent Nos. 4,803,279, 4,719,218 and 5,002,942, which references are hereby incorporated in their entirety. Suitable oxidising systems include sulfur in an inert solvent or solvent mixture, such as decalin, decalin and diglyme, p-cymene, xylene or mesitylene, under reflux conditions, or, preferably, potassium t-butoxide in t-butanol with dry air or oxygen.

15 Once the imidazole nucleus has been established, further compounds of formula (I) which may be prepared by applying standard techniques for functional group interconversion, for instance: -C(O)NR8R9 from -CO2CH3 by heating with or without catalytic metal cyanide, e.g. NaCN, and HNR8R9 in CH3OH; -OC(O)R8 from -OH with e.g., Cl(C)O)R8 in pyridine; -NR10-C(S)NR8R9 from -NHR10 with an alkylisothiocyanate or thiocyanate acid; NRC(O)OR6 from -NHR6 with the alkyl chloroformate; -NR10-C(O)NR8R9 from -NHR10 by treatment with an isocyanate, e.g. HN=O or R10N=C=O; -NR10-C(O)R8 from -NHR10 by treatment with Cl-C(O)R8 in pyridine; -C(-NR10)NR8R9 from -C(NR8R9)SR8 with HgNR8+OAc- by heating in alcohol; -C(NR8R9)SR8 from -C(S)NR8R9 with R8-I in an inert solvent, e.g. acetone; -C(S)NR8R9 (where R8 or R9 is not hydrogen) from -C(S)NH2 with HNR8R9, -C(-NCN)-NR8R9 from -C(-NR8R9)-SR8 with NH2CN by heating in anhydrous alcohol, alternatively from -C(-NH)-NR8R9 by treatment with BrCN and NaOEt in EtOH; -NR10-C(-NCN)SR8 from -NHR10 by treatment with (R8S)2C=NCN; -NR10SO2R8 from -NHR10 by treatment with ClSO2R8 by heating in pyridine; -NR10C(S)R8 from -NR10C(O)R8 by treatment with Lawesson's reagent [2,4-bis(4-methoxyphenyl)-1,3,2,4-dithiadiphosphetane-2,4-disulfide]; -NR10SO2CF3 from -NHR6 with triflic anhydride and base; -NR10C(O)-C(O)-OR8 from -NHR10 with, e.g. methoxymalyl chloride and a base such as triethylamine; -NR10C(O)-C(O)-NR8R9 from -NR10C(O)-C(O)-OR8 with HNR8R9; and 1-(NR10)-2-imidazolyl from -C(-NH)NR10 by heating with 2-chloroacetaldehyde in chloroform (wherein R6, R8, R9 and R10 are as hereinbefore defined).
Compounds of formula (I) in which R₂ is hydrogen may be readily converted into further compounds of formula (I) in which R₂ is other than hydrogen, for instance alkyl, by conventional procedures such as alkylation or acylation followed by reduction. Such methods are in general relatively inefficient as they lack regiospecificity and the desired N-1 product has to be separated from the mixture of N-1 and N-3 products, for instance by chromatography or fractional crystallisation.

Suitable protecting groups for use with hydroxyl groups and the imidazole nitrogen are well known in the art and described in many references, for instance, Protecting Groups in Organic Synthesis, Greene T W, Wiley-Interscience, New York, 1981. Suitable examples of hydroxyl protecting groups include silyl ethers, such as t-butylidimethyl or t-butyldiphenyl, and alkyl ethers, such as methyl connected by an alkyl chain of variable link, (CR₁₀R₂₀)n. Suitable examples of imidazole nitrogen protecting groups include tetrahydropyranyl.

Pharmaceutically acid addition salts of compounds of formula (I) may be obtained in known manner, for example by treatment thereof with an appropriate amount of acid in the presence of a suitable solvent.

METHODS OF TREATMENT

The compounds of Formula (I) or a pharmaceutically acceptable salt thereof can be used in the manufacture of a medicament for the prophylactic or therapeutic treatment of any disease state in a human, or other mammal, which is exacerbated or caused by excessive or unregulated cytokine production by such mammal's cell, such as but not limited to monocytes and/or macrophages.

Compounds of formula (I) are capable of inhibiting proinflammatory cytokines, such as IL-1, IL-6, IL-8 and TNF and are therefore of use in therapy. IL-1, IL-8 and TNF affect a wide variety of cells and tissues and these cytokines, as well as other leukocyte-derived cytokines, are important and critical inflammatory mediators of a wide variety of disease states and conditions. The inhibition of these pro-inflammatory cytokines is of benefit in controlling, reducing and alleviating many of these disease states.

Accordingly, the present invention provides a method of treating a cytokine-mediated disease which comprises administering an effective cytokine-interfering amount of a compound of formula (I) or a pharmaceutically acceptable salt thereof.
In particular, compounds of formula (I) or a pharmaceutically acceptable salt thereof are of use in the prophylaxis or therapy of any disease state in a human, or other mammal, which is exacerbated by or caused by excessive or unregulated IL-1, IL-8 or TNF production by such mammal's cell, such as, but not limited to, monocytes and/or macrophages.

Accordingly, in another aspect, this invention relates to a method of inhibiting the production of IL-1 in a mammal in need thereof which comprises administering to said mammal an effective amount of a compound of formula (I) or a pharmaceutically acceptable salt thereof.

There are many disease states in which excessive or unregulated IL-1 production is implicated in exaccerbating and/or causing the disease. These include rheumatoid arthritis, osteoarthritis, endotoxemia and/or toxic shock syndrome, other acute or chronic inflammatory disease states such as the inflammatory reaction induced by endotoxin or inflammatory bowel disease, tuberculosis, atherosclerosis, muscle degeneration, multiple sclerosis, cachexia, bone resorption, psoriatic arthritis, Reiter's syndrome, rheumatoid arthritis, gout, traumatic arthritis, rubella arthritis and acute synovitis. Recent evidence also links IL-1 activity to diabetes, pancreatic β cells and Alzheimer's disease.

In a further aspect, this invention relates to a method of inhibiting the production of TNF in a mammal in need thereof which comprises administering to said mammal an effective amount of a compound of formula (I) or a pharmaceutically acceptable salt thereof.

Excessive or unregulated TNF production has been implicated in mediating or exacerbating a number of diseases including rheumatoid arthritis, rheumatoid spondylitis, osteoarthritis, gouty arthritis and other arthritic conditions, sepsis, septic shock, endotoxic shock, gram negative sepsis, toxic shock syndrome, adult respiratory distress syndrome, cerebral malaria, chronic pulmonary inflammatory disease, silicosis, pulmonary sarcoidosis, bone resorption diseases, such as osteoporosis, reperfusion injury, graft vs. host reaction, allograft rejections, fever and myalgias due to infection, such as influenza, cachexia secondary to infection or malignancy, cachexia secondary to acquired immune deficiency syndrome (AIDS), AIDS, ARC (AIDS related complex), keloid formation, scar tissue formation, Crohn's disease, ulcerative colitis and pyresis.

Compounds of formula (I) are also useful in the treatment of viral infections, where such viruses are sensitive to upregulation by TNF or will elicit TNF production in vivo. The viruses contemplated for treatment herein are those that produce TNF as a result of infection, or those which

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are sensitive to inhibition, such as by decreased replication, directly or indirectly, by the TNF inhibiting-compounds of formula (1). Such viruses include, but are not limited to HIV-1, HIV-2 and HIV-3, Cytomegalovirus (CMV), Influenza, adenovirus and the Herpes group of viruses, such as but not limited to, Herpes Zoster and Herpes Simplex. Accordingly, in a further aspect, this invention relates to a method of treating a mammal afflicted with a human immunodeficiency virus (HIV) which comprises administering to such mammal an effective TNF inhibiting amount of a compound of formula (I) or a pharmaceutically acceptable salt thereof.

Compounds of formula (I) may also be used in association with the veterinary treatment of mammals, other than in humans, in need of inhibition of TNF production. TNF mediated diseases for treatment, therapeutically or prophylactically, in animals include disease states such as those noted above, but in particular viral infections. Examples of such viruses include, but are not limited to, the lentivirus infections such as equine infectious anaemia virus, caprine arthritis virus, visna virus, or the maedi virus, or the retroviruses, such as feline immunodeficiency virus (FIV), bovine immunodeficiency virus, or canine immunodeficiency virus.

The compounds of formula (I) may also be used topically in the treatment or prophylaxis of topical disease states mediated by or exacerbated by excessive cytokine production, such as by IL-1 or TNF respectively, such as inflamed joints, eczema, psoriasis and other inflammatory skin conditions such as sunburn; inflammatory eye conditions including conjunctivitis; pyresis, pain and other conditions associated with inflammation.

Compounds of formula (I) have also been shown to inhibit the production of IL-8 (Interleukin-8, NAP). Accordingly, in a further aspect, this invention relates to a method of inhibiting the production of IL-8 in a mammal in need thereof which comprises administering to said mammal an effective amount of a compound of formula (I) or a pharmaceutically acceptable salt thereof.

There are many disease states in which excessive or unregulated IL-8 production is implicated in exacerbating and/or causing the disease. These diseases are characterized by massive neutrophil infiltration such as, psoriasis, inflammatory bowel disease, asthma, cardiac and renal reperfusion injury, adult respiratory distress syndrome, thrombosis and glomerulonephritis. All of these diseases are associated with increased IL-8 production which is responsible for the chemotaxis of neutrophils into the inflammatory site. In contrast to other inflammatory cytokines (IL-1, TNF,
and IL-6), IL-8 has the unique property of promoting neutrophil chemotaxis and activation. Therefore, the inhibition of IL-8 production would lead to a direct reduction in the neutrophil infiltration.

The compounds of formula (I) are administered in an amount sufficient to inhibit cytokine, in particular IL-1, IL-8 or TNF, production such that it is regulated down to normal levels, or in some case to subnormal levels, so as to ameliorate or prevent the disease state. Abnormal levels of IL-1, IL-8 or TNF, for instance in the context of the present invention, constitute: (i) levels of free (not cell bound) IL-1, IL-8 or TNF greater than or equal to 1 picogram per ml; (ii) any cell associated IL-1, IL-8 or TNF; or (iii) the presence of IL-1, IL-8 or TNF mRNA above basal levels in cells or tissues in which IL-1, IL-8 or TNF, respectively, is produced.

The discovery that the compounds of formula (I) are inhibitors of cytokines, specifically IL-1, IL-8 and TNF is based upon the effects of the compounds of formulas (I) on the production of the IL-1, IL-8 and TNF in vitro assays which are described herein.

As used herein, the term "inhibiting the production of IL-1 (IL-8 or TNF)" refers to:

a) a decrease of excessive in vivo levels of the cytokine (IL-1, IL-8 or TNF) in a human to normal or sub-normal levels by inhibition of the in vivo release of the cytokine by all cells, including but not limited to monocytes or macrophages;

b) a down regulation, at the genomic level, of excessive in vivo levels of the cytokine (IL-1, IL-8 or TNF) in a human to normal or sub-normal levels;

c) a down regulation, by inhibition of the direct synthesis of the cytokine (IL-1, IL-8 or TNF) as a posttranslational event; or

d) a down regulation, at the translational level, of excessive in vivo levels of the cytokine (IL-1, IL-8 or TNF) in a human to normal or sub-normal levels.

As used herein, the term "TNF mediated disease or disease state" refers to any and all disease states in which TNF plays a role, either by production of TNF itself, or by TNF causing another monokine to be released, such as but not limited to IL-1, IL-6 or IL-8. A disease state in which, for instance, IL-1 is a major component, and whose production or action, is exacerbated or secreted in response to TNF, would therefore be considered a disease stated mediated by TNF.
As used herein, the term "cytokine" refers to any secreted polypeptide that affects the functions of cells and is a molecule which modulates interactions between cells in the immune, inflammatory or hematopoietic response. A cytokine includes, but is not limited to, monokines and lymphokines, regardless of which cells produce them. For instance, a monokine is generally referred to as being produced and secreted by a mononuclear cell, such as a macrophage and/or monocyte. Many other cells however also produce monokines, such as natural killer cells, fibroblasts, basophils, neutrophils, endothelial cells, brain astrocytes, bone marrow stromal cells, epideral keratinocytes and B-lymphocytes. Lymphokines are generally referred to as being produced by lymphocyte cells. Examples of cytokines include, but are not limited to, Interleukin-1 (IL-1), Interleukin-6 (IL-6), Interleukin-8 (IL-8), Tumor Necrosis Factor-alpha (TNF-α) and Tumor Necrosis Factor beta (TNF-β).

As used herein, the term "cytokine interfering" or "cytokine suppressive amount" refers to an effective amount of a compound of formula (I) which will cause a decrease in the in vivo levels of the cytokine to normal or sub-normal levels, when given to a patient for the prophylaxis or treatment of a disease state which is exacerbated by, or caused by, excessive or unregulated cytokine production.

As used herein, the cytokine referred to in the phrase "inhibition of a cytokine, for use in the treatment of a HIV-infected human" is a cytokine which is implicated in (a) the initiation and/or maintenance of T cell activation and/or activated T cell-mediated HIV gene expression and/or replication and/or (b) any cytokine-mediated disease associated problem such as cachexia or muscle degeneration.

As TNF-β (also known as lymphotoxin) has close structural homology with TNF-α (also known as cachectin) and since each induces similar biologic responses and binds to the same cellular receptor, both TNF-α and TNF-β are inhibited by the compounds of the present invention and thus are herein referred to collectively as "TNF" unless specifically delineated otherwise.

In order to use a compound of formula (I) or a pharmaceutically acceptable salt thereof in therapy, it will normally be formulated into a pharmaceutical composition in accordance with standard pharmaceutical practice. This invention, therefore, also relates to a pharmaceutical composition comprising an effective, non-toxic amount of a compound of formula (I) and a pharmaceutically acceptable carrier or diluent.
Compounds of formula (I), pharmaceutically acceptable salts thereof and pharmaceutical compositions incorporating such may conveniently be administered by any of the routes conventionally used for drug administration, for instance, orally, topically, parenterally or by inhalation. The compounds of formula (I) may be administered in conventional dosage forms prepared by combining a compound of formula (I) with standard pharmaceutical carriers according to conventional procedures. The compounds of formula (I) may also be administered in conventional dosages in combination with a known, second therapeutically active compound. These procedures may involve mixing, granulating and compressing or dissolving the ingredients as appropriate to the desired preparation. It will be appreciated that the form and character of the pharmaceutically acceptable character or diluent is dictated by the amount of active ingredient with which it is to be combined, the route of administration and other well-known variables. The carrier(s) must be "acceptable" in the sense of being compatible with the other ingredients of the formulation and not deleterious to the recipient thereof.

The pharmaceutical carrier employed may be, for example, either a solid or liquid. Exemplary of solid carriers are lactose, terra alba, sucrose, talc, gelatin, agar, pectin, acacia, magnesium stearate, stearic acid and the like. Exemplary of liquid carriers are syrup, peanut oil, olive oil, water and the like. Similarly, the carrier or diluent may include time delay material well known to the art, such as glyceryl mono-stearate or glyceryl distearate alone or with a wax.

A wide variety of pharmaceutical forms can be employed. Thus, if a solid carrier is used, the preparation can be tableted, placed in a hard gelatin capsule in powder or pellet form or in the form of a troche or lozenge. The amount of solid carrier will vary widely but preferably will be from about 25mg. to about 1g. When a liquid carrier is used, the preparation will be in the form of a syrup, emulsion, soft gelatin capsule, sterile injectable liquid such as an ampule or nonaqueous liquid suspension.

Compounds of formula (I) may be administered topically, that is by non-systemic administration. This includes the application of a compound of formula (I) externally to the epidermis or the buccal cavity and the instillation of such a compound into the ear, eye and nose, such that the compound does not significantly enter the blood stream. In contrast, systemic administration refers to oral, intravenous, intraperitoneal and intramuscular administration.
Formulations suitable for topical administration include liquid or semi-liquid preparations suitable for penetration through the skin to the site of inflammation such as liniments, lotions, creams, ointments or pastes, and drops suitable for administration to the eye, ear or nose. The active ingredient may comprise, for topical administration, from 0.001% to 10% w/w, for instance from 1% to 2% by weight of the formulation. It may however comprise as much as 10% w/w but preferably will comprise less than 5% w/w, more preferably from 0.1% to 1% w/w of the formulation.

Lotions according to the present invention include those suitable for application to the skin or eye. An eye lotion may comprise a sterile aqueous solution optionally containing a bactericide and may be prepared by methods similar to those for the preparation of drops. Lotions or liniments for application to the skin may also include an agent to hasten drying and to cool the skin, such as an alcohol or acetone, and/or a moisturizer such as glycerol or an oil such as castor oil or arachis oil.

Creams, ointments or pastes according to the present invention are semi-solid formulations of the active ingredient for external application. They may be made by mixing the active ingredient in finely-divided or powdered form, alone or in solution or suspension in an aqueous or non-aqueous fluid, with the aid of suitable machinery, with a greasy or non-greasy base. The base may comprise hydrocarbons such as hard, soft or liquid paraffin, glycerol, beeswax, a metallic soap; a mucilage; an oil of natural origin such as almond, corn, arachis, castor or olive oil; wool fat or its derivatives or a fatty acid such as steric or oleic acid together with an alcohol such as propylene glycol or a macrogel. The formulation may incorporate any suitable surface active agent such as an anionic, cationic or non-ionic surfactant such as a sorbitan ester or a polyoxyethylene derivative thereof. Suspending agents such as natural gums, cellulose derivatives or inorganic materials such as siliceous silicas, and other ingredients such as lanolin, may also be included.

Drops according to the present invention may comprise sterile aqueous or oily solutions or suspensions and may be prepared by dissolving the active ingredient in a suitable aqueous solution of a bactericidal and/or fungicidal agent and/or any other suitable preservative, and preferably including a surface active agent. The resulting solution may then be clarified by filtration, transferred to a suitable container which is then sealed and sterilized by autoclaving or maintaining at 98-100°C. for half an hour. Alternatively, the solution may be sterilized by filtration and transferred to the container by an aseptic technique. Examples of
bactericidal and fungicidal agents suitable for inclusion in the drops are 
phenylmercuric nitrate or acetate (0.002%), benzalkonium chloride (0.01%) 
and chlorhexidine acetate (0.01%). Suitable solvents for the preparation of 
an oily solution include glycerol, diluted alcohol and propylene glycol.

Compounds of formula (I) may be administered parenterally, that is 
by intravenous, intramuscular, subcutaneous intranasal, intrarectal, 
intravaginal or intraperitoneal administration. The subcutaneous and 
intramuscular forms of parenteral administration are generally preferred. 
Appropriate dosage forms for such administration may be prepared by 
conventional techniques. Compounds of formula (I) may also be 
administered by inhalation, that is by intranasal and oral inhalation 
administration. Appropriate dosage forms for such administration, such as 
an aerosol formulation or a metered dose inhaler, may be prepared by 
conventional techniques.

For all methods of use disclosed herein for the compounds of formula 
(I), the daily oral dosage regimen will preferably be from about 0.1 to about 
80 mg/kg of total body weight, preferably from about 0.2 to 30 mg/kg, more 
preferably from about 0.5 mg to 15 mg. The daily parenteral dosage regimen 
about 0.1 to about 80 mg/kg of total body weight, preferably from about 0.2 
to about 30 mg/kg, and more preferably from about 0.5 mg to 15 mg/kg. The 
daily topical dosage regimen will preferably be from 0.1 mg to 150 mg, 
administered one to four, preferably two or three times daily. The daily 
inhalation dosage regimen will preferably be from about 0.01 mg/kg to about 
1 mg/kg per day. It will also be recognized by one of skill in the art that the 
optimal quantity and spacing of individual dosages of a compound of 
formula (I) or a pharmaceutically acceptable salt thereof will be determined 
by the nature and extent of the condition being treated, the form, route and 
site of administration, and the particular patient being treated, and that 
such optimums can be determined by conventional techniques. It will also 
be appreciated by one of skill in the art that the optimal course of 
treatment, i.e., the number of doses of a compound of formula (I) or a 
pharmaceutically acceptable salt thereof given per day for a defined number 
of days, can be ascertained by those skilled in the art using conventional 
course of treatment determination tests.

The invention will now be described by reference to the following 
examples which are merely illustrative and are not to be construed as a 
limitation of the scope of the present invention.
Synthetic Examples

Example 1 - 2-(4-Cyanophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - To a solution of 2-(4-cyanophenyl)-4-(4-fluorophenyl)-N-1-hydroxy-5-(4-pyridyl)imidazole (4.5 g, 13.2 mmol) [See Ex. 10 below] in DMF (50 mL) was added triethyl phosphite (3.4 mL, 20 mmol), and the resulting mixture was heated at 100 °C for 2 h. After cooling, the mixture was poured into H₂O, and the solid which formed was collected by filtration, washed with H₂O and dried in vacuo to afford the title compound (4.0 g, 89%). Recrystallization from CH₂Cl₂/MeOH gave white solid with a mp of 268-269 °C.

Example 2 - 1-Methyl-2-(4-methoxyphenyl)-4-phenyl-5-(4-pyridyl)-imidazole - (a) N-Methyl-N-(4-picolyl)amine- To 4-picolyl chloride, hydrochloride (10 g, 0.06 mol) was added methylamine (50 mL of 40% aqueous solution, 0.58 mol), and the resulting purple solution was stirred at rt for 30 min, then poured into H₂O. The mixture was extracted with CH₂Cl₂ (6x), and the combined organic extracts were evaporated. The residue was filtered under reduced pressure through a silica gel column, eluting with a solvent gradient of 0-10% MeOH/CHCl₃ to provide the title compound as a light yellow oil (4.8 g, 66%): ¹H NMR (CDCl₃): δ 8.50 (dd, 2H); 7.20 (dd, 2H); 3.70 (s, 2H); 2.40 (s, 3H); 1.70 (br, 1H).

(b) 4-Methoxy-N-methyl-N-(4-picolyl)benzamide - To a solution of N-methyl-N-(4-picolyl)amine (0.40 g, 3.3 mmol) and triethylamine (1.5 mL, 10.8 mmol) in CH₂Cl₂ (15 mL) was added 4-methoxybenzoyl chloride (1.2 g, 7.3 mmol). The resulting mixture was stirred at rt for 15 min, and then partitioned between 2.5N NaOH and Et₂O. The organic extract was washed with saturated aqueous NaCl and dried (MgSO₄). The solvent was removed in vacuo, and the residue was purified by flash chromatography, eluting with a solvent gradient of 2-4% MeOH/CHCl₃. The material that was isolated was triturated with Et₂O to provide the title compound as a light yellow solid (0.18 g, 21%): ¹H NMR (CDCl₃): δ 8.60 (d, 2H); 7.43 (br d, 2H); 7.20 (br s, 2H); 6.90 (br d, 2H); 4.66 (br s, 2H); 3.80 (s, 3H); 3.00 (s, 3H).

(c) 1-Methyl-2-(4-methoxyphenyl)-4-phenyl-5-(4-pyridyl)-imidazole - To a solution of diisopropylamine (0.16 mL, 1.1 mmol) in THF at -78 °C was added n-butyllithium (0.38 mL of 2.5 M solution, 0.95 mmol). To the resulting mixture was added a solution of 4-methoxy-N-methyl-N-(4-picolyl)benzamide (0.16 g, 0.62 mmol) in THF. The resulting dark red solution was warmed to -40 °C and stirred for 15 min, at which time benzonitrile (0.13 mL, 1.2 mmol) was added. The mixture was warmed to rt
and stirred for 10 h. Aqueous NH$_4$Cl (0.5 mL) was added, and the mixture was concentrated under reduced pressure. The residue was purified by flash chromatography, eluting with a solvent gradient of 2-4% MeOH/CHCl$_3$. The material which was isolated was triturated with Et$_2$O and recrystallized from EtOAc to provide the title compound as an off-white solid (35 mg, 17%): mp 193-194 ºC.

Example 3 - 2-(4-Cyanophenyl)-1-methyl-4-phenyl-5-(4-pyridyl)imidazole - (a) 4-Cyano-N-methyl-N-(4-picoly)benzamide - The title compound was prepared using the same procedure as described in Example 2, step (b) except using 4-cyanobenzoyl chloride: $^1$H NMR (CDCl$_3$): $\delta$ 8.49 (dd, 2H); 7.86-7.04 (m, 6H); 4.70 and 4.43 (two br s, 2H); 3.08 and 2.89 (two br s, 3H).

(b) 4-Cyano-N-[N'-α-dibenzoyl-1,4-dihydropyridylmethylenyl]-N-methylbenzamide - To a solution of diisopropylamine (2.8 mL, 20 mmol) in THF at -78 ºC was added n-butyllithium (6.7 mL of 2.5 M solution, 17 mmol). To the resulting mixture was added a solution of 4-cyano-N-methyl-N-(4-picoly)benzamide (3.5 g, 14 mmol) in THF. The resulting dark purple solution was stirred at -78 ºC for 10 min, at which time benzyol chloride (4.1 mL, 35 mmol) was added. The mixture was warmed to room temperature over 30 min, then poured into aqueous NH$_4$Cl. The mixture was extracted with Et$_2$O, and the organic extract was evaporated under reduced pressure. The residue was triturated with Et$_2$O to provide an orange solid which was washed sparingly with acetone and copiously with Et$_2$O. The title compound was obtained as a bright yellow solid (1.6 g, 25%): $^1$H NMR (CDCl$_3$): $\delta$ 7.81-7.09 (m, 16H); 6.49 (m, 2H); 3.32 (s, 3H).

(c) 2-(4-Cyanophenyl)-1-methyl-4-phenyl-5-(4-pyridyl)imidazole - To a solution of 4-cyano-N-[N'-α-dibenzoyl-1,4-dihydropyridylmethylenyl]-N-methylbenzamide (1.5 g, 3.3 mmol) in acetic acid (50 mL) was added ammonium acetate (1.5 g, 19.5 mmol). The resulting mixture was heated at reflux for 18 h, then allowed to cool and was concentrated. The residue was suspended in CH$_2$Cl$_2$ and filtered. The filtrate was evaporated and the residue was triturated with MeOH to afford the title compound as a white crystalline solid (0.72 g, 64%): mp 176-177 ºC.

Example 4 - 2-(4-Aminomethylphenyl)-1-methyl-4-phenyl-5-(4-pyridyl)-imidazole - To a solution of 2-(4-cyanophenyl)-1-methyl-4-phenyl-5-(4-pyridyl)imidazole (0.20 g, 0.6 mmol) [See Ex. 3 above] in THF (10 mL) was added LiAlH$_4$ (0.60 mL of 1.0 M solution in THF, 0.6 mmol), and the
resulting mixture was stirred at rt for 1 h. The mixture was then poured into 2.5 N NaOH and extracted with Et₂O. The organic extract was evaporated, and the residue was purified by flash chromatography, eluting first with a solvent gradient of 0-10% MeOH/CHCl₃, followed by 1:10:90 NH₄OH/MeOH/CHCl₃. Trituration with ether afforded the title compound as a white solid (66 mg, 32%): CIMS (NH₃, m/z): 341 (M⁺+H).  

**Example 5** - 4-[1-Methyl-4-phenyl-5(4-pyridyl)-imidazol-2-yl] benzoic acid, sodium salt - A mixture of 2-(4-cyanophenyl)-1-methyl-4-phenyl-5(4-pyridyl)imidazole (0.10 g, 0.3 mmol) [See Ex. 3 above] in 6 N HCl (3 mL) was heated at reflux for 24 h, then allowed to cool. The pH was adjusted to 7, and the solid which formed was collected by filtration and washed successively with H₂O, acetone and Et₂O to provide the title compound as a white solid (25 mg, 23%): CIMS (NH₃, m/z): 356 (M⁺+H).  

**Example 6** - 2-(4-Acetamidomethylenephynyl)-1-methyl-4-phenyl-5-(4-pyridyl)imidazole - To a solution of 2-(4-aminomethylphenyl)-1-methyl-4-phenyl-5-(4-pyridyl)imidazole (30 mg, 0.09 mmol) [See Ex. 4 above] in pyridine (3 mL) was added acetic anhydride (0.30 mL, 3.18 mmol). The resulting solution was stirred at rt for 30 min, then concentrated under reduced pressure. The residue was purified by flash chromatography, eluting with a solvent gradient of 0-2% MeOH/CHCl₃. The isolated material was triturated with Et₂O to provide an off-white solid (10 mg, 28%) which was recrystallized from EtOAc to provide the title compound: mp 210-211 °C.  

**Example 7** - Methyl-4-[1-methyl-4-phenyl-5-(4-pyridyl)-imidazol-2-yl] benzoate - To a suspension of 4-[1-methyl-4-phenyl-5(4-pyridyl)-imidazol-2-yl] benzoic acid, sodium salt (20 mg, 0.06 mmol) [See Ex. 5 above] in CH₂Cl₂ (2 mL) was added triethylamine (24 mL, 0.17 mmol), followed by thionyl chloride (10 mL, 0.14 mmol). The reaction mixture was stirred at rt for 30 min, at which time MeOH (0.5 mL) was added. The mixture was stirred at rt for an additional 2 h and concentrated under reduced pressure. The residue was purified by flash chromatography, eluting with a solvent gradient of 0-1% MeOH/CHCl₃ and recrystallized from EtOAc to afford the title compound as an off-white crystalline solid (1.6 mg, 8%): mp 208-209 °C.  

**Example 8a** - 4-(4-Fluorophenyl)-N-1-hydroxy-2-(4-hydroxyphenyl)-5-(4-pyridyl)imidazole - The title compound was prepared using the same procedure as described in Example 10, step (d) except using 4-hydroxybenzaldehyde.  

**Example 8b** - 4-(4-Fluorophenyl)-2-(4-hydroxyphenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure.
as described in Example 1, except using 4-(4-fluorophenyl)-N-1-hydroxy-2-
(4-hydroxyphenyl)-5-(4-pyridyl)imidazole (see Ex.8a): mp 214-215 °C.

**Example 9** - 4-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-
yl]benzoic acid - A solution containing 2-(4-cyanophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole (9.6 g, 28 mmol) [See Ex. 1 above] in concentrated HCl (100 mL) was heated at reflux for 18 h. After cooling, the pH was adjusted to neutral by the addition of 50% aqueous NaOH. The solid which formed was collected by filtration and washed successively with H₂O, acetone and Et₂O. A portion of the solid (5 g) was dissolved in MeOH and filtered under reduced pressure through a pad of silica gel, eluting with a solvent gradient of 4-10 % MeOH/CHCl₃, followed by 2:20:80 H₂O/MeOH/CHCl₃. The title compound was isolated as a yellow solid, which was re-crystallized from MeOH/CH₂Cl₂ (1.2 g, 30% adjusted yield): mp 289-290 °C.

**Example 10** - 2-(4-Cyanophenyl)-4-(4-fluorophenyl)-1-N-hydroxy-5-(4-
pyridyl)imidazole - (a) 4-Fluoro-N-methoxy-N-methylbenzamidine
- To a mixture containing methoxymethylamine hydrochloride (4 g, 0.45 mol) and triethylamine (138 mL, 0.99 mol) in CH₂Cl₂ (500 mL) at 0 °C was added over 30 min, 4-fluorobenzoyl chloride (50 mL, 0.41 mol). The resulting mixture was allowed to warm to rt and stirring was continued for 30 min, at which time the mixture was poured into H₂O and extracted with Et₂O. The organic extract was washed with saturated aqueous NaCl and dried (MgSO₄). Removal of the solvent *in vacuo* afforded the title compound (80 g, 100%), which was used without further purification: ¹H NMR (CDCl₃): δ 7.72 (dd, 2H); 7.06 (apparent t, 2H); 3.52 (s, 3H); 3.43 (s, 3H).

(b) 4-Fluoro-2-(4-pyridyl)acetophenone - A solution of lithium diisopropylamide was prepared at -78 °C in the usual manner from diisopropylamine (21 ml, 0.15 mol) and n-butyllithium (54 mL of 2.5 M solution in hexanes, 0.135 mol), and to this was added at -78 °C, 4-picoline (10 g, 0.108 mol). After stirring an additional 15 min at -78 °C, 4-fluoro-N-
methoxy-N-methylbenzamidine (20 g, 0.109 mol) was added, and the mixture was allowed to slowly warm to rt. The reaction mixture was poured into saturated aqueous NaCl and extracted with 4:1 THF/CH₂Cl₂, and the organic extract was dried (MgSO₄). The solvent was removed *in vacuo*, and to the oily brown residue was added Et₂O. The title compound was obtained as a brown solid (16.8 g, 72%) which was recrystallized from Et₂O/Hex: ¹H NMR (CDCl₃): δ 8.55 (d, 2H); 8.03 (dd, 2H); 7.16 (m, 4H); 4.24 (s, 2H).

(c) 4-Fluoro-2-hydroxyimino-2-(4-pyridyl)acetophenone - The title compound was prepared using the same procedure (US 3,940,486)
employed to prepare 2-hydroxyimino-2-(4-pyridyl)acetophenone, except using 4-fluoro-2-(4-pyridyl)acetophenone.

(d) 2-(4-Cyanophenyl)-4-(4-fluorophenyl)-N-1-hydroxy-5-(4-pyridyl)imidazole - The title compound was prepared using the same procedure (US 3,940,486) employed to prepare 2-(t-butyl)-4-(phenyl)-N-1-hydroxy-5-(4-pyridyl)imidazole, except using 4-fluoro-2-hydroxyimino-2-(4-pyridyl)acetophenone and 4-cyanobenzaldehyde: $^1$H NMR (CDCl$_3$): $\delta$ 8.27 (d, 2H); 7.94 (d, 2H); 7.72 (d, 2H); 7.35 (d, 2H); 7.30 (dd, 2H); 6.96 (t, 2H).

Example 11 - 2-(4-Aminomethylphenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - To a solution of 2-(4-cyanophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole (2.5 g, 7.3 mmol) [See Ex. 1 above] in THF (50 mL) was added LiAlH$_4$ (7.3 mL of 1 M solution in THF, 7.3 mmol), and the resulting mixture was heated at reflux for 2 h, at which time tlc analysis indicated that the reaction was incomplete. Additional LiAlH$_4$ (4.0 mL, 4.0 mmol) was added and heating was continued for 30 min. The mixture was allowed to cool, then poured into 2.5 N NaOH and extracted with THF. The organic extract was washed with saturated aqueous NaCl and concentrated under reduced pressure. The residue was purified by flash chromatography, eluting with 9:1 CHCl$_3$/MeOH, followed by 90:10:1 CHCl$_3$/MeOH/NH$_3$. The material that was isolated was triturated with Et$_2$O to afford the title compound (1.5 g, 60%): mp 214-215 °C.

Example 12a - 2-(4-Cyanophenyl)-4-(4-fluorophenyl)-N-1-hydroxy-5-(4-quinolyl)imidazole - (a) 4-Fluoro-2-(4-quinolyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (b) except using 4-methylquinoline: $^1$H NMR (CDCl$_3$): $\delta$ 8.87 (d, 1H); 8.13 (m, 3H); 7.86 (d, 1H); 7.73 (apparent br t, 1H); 7.56 (apparent br t, 1H); 7.28 (d, 1H); 7.20 (t, 2H); 4.71 (s, 2H).

(b) 4-Fluoro-2-hydroxyimino-2-(4-quinolyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (c) except using 4-fluoro-2-(4-quinolyl)acetophenone: $^1$H NMR (DMSO-d$_6$): $\delta$ 9.00 (d, 1H); 8.15 (m, 3H); 7.78 (m, 1H); 7.61 (m, 2H); 7.50 (d, 1H); 7.42 (t, 2H).

(c) 2-(4-Cyanophenyl)-4-(4-fluorophenyl)-N-1-hydroxy-5-(4-quinolyl)imidazole - The title compound was prepared using the same procedure as described in Example 10, step (d) except using 4-fluoro-2-hydroxyimino-2-(4-quinolyl)acetophenone and 4-cyanobenzaldehyde: $^1$H NMR (CDCl$_3$): $\delta$ 8.30 (d, 2H); 7.80 (d, 1H); 7.70 (two overlapping d, 3H); 7.46 (m, 2H); 7.36 (m, 1H); 7.11 (m, 2H); 7.01 (m, 1H); 6.75 (t, 2H).
Example 12b - 2-(4-Cyanophenyl)-4-(4-fluorophenyl)-5-(4-quinolyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 1, except using 2-(4-cyanophenyl)-4-(4-fluorophenyl)-N-1-hydroxy-5-(4-quinolyl)imidazole [see Ex. 12a]: mp 294-295 °C.

Example 13 - 2-(3,5-Dibromo-4-hydroxyphenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - (a) 1-(4-fluorophenyl)-2-(4-pyridyl)-ethanediol To a stirring solution of 2.0 g (11.2 mmol) 4-(t-butylidimethylsilyloxy)methyl pyridine in 8 ml of THF at -20 °C was added 14.7 mmol of lithium di-iso-propyl amide in THF. Thirty minutes later 4-fluorobenzaldehyde (1.66 g, 13.4 mmol) was added at which point the solution was allowed to warm slowly to rt. The reaction was quenched with NH₄Cl and extracted with ether to afford the crude protected diol which following concentration was dissolved in THF and treated with 17 ml of a 1 molar solution of tetrabutylammonium fluoride in THF overnight. Standard aqueous workup afforded the crude diol which was further purified by column chromatography (hex/EtOAc) to yield 1.6 g (62%) of the titled material.

(b) 1-(4-fluorophenyl)-2-(4-pyridyl)ethanedione Oxidation of 1-(4-fluorophenyl)-2-(4-pyridyl)ethanediol according to the oxalyl chloride method of Swern [J. Org. Chem., 44, p 4148, 1979)] gave the titled dione following extractive workup and recrystallization from hexanes m.p. 85-86.5 °C.

(c) 2-(3,5-Dibromo-4-hydroxyphenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole To a solution of 1-(4-fluorophenyl)-2-(4-pyridyl)ethanediol (0.25 g, 1.1 mmol) and 3,5-dibromo-4-hydroxybenzaldehyde (0.37 g, 1.3 mmol) in glacial acetic acid (5 mL) was added ammonium acetate (0.50 g, 6.5 mmol), and the resulting mixture was heated at reflux for 18 h. After cooling, the mixture was poured into H₂O, and the pH was adjusted to neutral by the addition of 2.5 N NaOH. The solid which formed was collected by filtration, washed with H₂O, dried in vacuo and purified by flash chromatography, eluting with a solvent gradient of 2-4% MeOH/CHCl₃. The title compound was obtained as a tan solid (15 mg, 3%): ESI-MS (m/z): 488 (M+H).
white solid (3.2 g 66%). 1H NMR (CDCl3/MeOH-d4): δ 8.45 (d, 2H); 8.12 (m, 4H); 7.52 (m, 4H); 7.15 (t, 2H); 4.42 (q, 2H); 1.43 (t, 3H).

**Example 15 - 2-[3,5-Dimethyl-4-hydroxy(phenyl)]-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole** - The title compound was prepared using the same procedure as described in Example 13, except using 3,5-dimethyl-4-hydroxybenzaldehyde: ESMS (m/z): 360 (M+H).

**Example 16 - 4-(4-Fluorophenyl)-2-(2-hydroxyphenyl)-5-(4-pyridyl)-1H-imidazole** - The title compound was prepared using the same procedure as described in Example 13, except using salicylaldehyde: ESMS (m/z): 332 (M+H).

**Example 17 - 4-(4-Fluorophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole** - The title compound was prepared using the same procedure as described in Example 13, except using 4-(methylthio)benzaldehyde: ESMS (m/z): 362 (M+H).

**Example 18 - Methyl 4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]-benzoate** - A mixture containing 4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzoic acid, sodium salt (0.20 g, 0.5 mmol) [See Ex. 9 above] and concentrated HCl (10 drops) in MeOH (5 mL) was heated at reflux for 8 h. After cooling, the pH was adjusted to neutral by the addition of 2.5 N NaOH, and the solid which formed was collected by filtration, washed with H2O and dried in vacuo. The title compound was obtained as a yellow solid (0.14 g, 76%) and was recrystallized from EtOAc/CH2Cl2: 1H NMR (CDCl3/MeOH-d4): δ 8.36 (d, 2H); 8.03 (m, 4H); 7.60-7.30 (m, 4H); 7.07 (t, 2H); 3.84 (s,3H).

**Example 19 - 4-(4-Fluorophenyl)-2-(4-methylsulfonylphenyl)-5-(4-pyridyl)-1H-imidazole** - To a solution of 4-(4-fluorophenyl)-2-(4-methylsulfonylphenyl)-5-(4-pyridyl)-1H-imidazole (3.7 g, 9.8 mmol) [See Ex. 20 below] in 1:10 3 N HCl/H2O (88 mL) was added a solution of KMnO4 (1.5 g, 9.8 mmol) in H2O (15 mL). After stirring at rt for 1 h, additional KMnO4 (0.15 g, 0.9 mmol) was added, and stirring was continued for 15 min. The mixture was then poured into saturated aqueous Na2SO3 (200 mL), and the pH was adjusted to slightly acidic by the addition of 3 N HCl, then to neutral by the addition of 2.5 N NaOH. The solid which formed was collected by filtration, washed successively with H2O and MeOH and recrystallized three times from MeOH to afford the title compound (0.63 g, 16%); mp 148-149 °C.

**Example 20 - 4-(4-Fluorophenyl)-2-(4-methylsulfinylphenyl)-5-(4-pyridyl)-1H-imidazole** - To a solution of 4-(4-fluorophenyl)-2-(4-
methylthiophenyl)-5-(4-pyridyl)-1H-imidazole (0.80 g, 2.2 mmol) [See Ex. 17 above] in glacial acetic acid (15 mL) was added a solution of K₂S₂O₅ (0.72 g, 2.6 mmol) in H₂O (20 mL). Additional glacial acetic acid (15 mL) was added to ensure homogeneity, and the resulting solution was stirred at rt for 18 h. The mixture was then poured into H₂O, and the pH was adjusted to neutral by the addition of conc. NH₄OH. The solid which formed was collected by filtration to afford the title compound (0.65 g, 78%) as a tan solid, which was recrystallized from MeOH: mp 250-252 °C.

**Example 21** - N,N-Dimethyl-4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzamide - To dimethylamino(methyl)aluminum chloride (0.60 mL of 0.67 M solution in toluene, 0.40 mmol) was added a solution of methyl 4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]-benzoate (50 mg, 0.13 mmol) [See Ex. 18 above] in 1,2-dichloroethane (5 mL). The resulting mixture was heated at reflux for 4 h, then allowed to cool and was poured into 2.5 N NaOH. The mixture was extracted with EtOAc, and the organic extract was washed with saturated aqueous NaCl and dried (MgSO₄). The solvent was removed in vacuo, and the residue was purified by flash chromatography, eluting with a solvent gradient of 2-4% MeOH/CHCl₃ to afford the title compound (25 mg, 50%) as a white solid: CIMS (NH₃, m/z): 387 (M+H).

**Example 22** - 2-[(4-N,N-Dimethylaminomethylphenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 11, except using N,N-dimethyl-4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzamide: CIMS (NH₃, m/z): 373 (M+H).

**Example 23** - 2-[4-(Dimethylamino)phenyl]-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 13, except using 4-(N,N-dimethylamino)benzaldehyde: ESMS (m/z): 359 (M+H).

**Example 24** - 4-(4-Fluorophenyl)-2-phenyl-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 13, except using benzaldehyde: ESMS (m/z): 316 (M+H).

**Example 25** - 2-[4-(3-Dimethylaminopropoxy)phenyl]-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 13, except using 4-(3-dimethylamino-propoxy)benzaldehyde: ESMS (m/z): 417 (M+H).

**Example 26** - 4-(4-Fluorophenyl)-2-(4-nitrophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as
described in Example 13, except using 4-nitrobenzaldehyde: EIMS (m/z): 359 (M+H).

**Example 27 - N,N-Dimethyl-4-[2-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzoyl-oxyacetamide -**

(a) Methyl benzylglycolate -

To a solution containing methyl glycolate (2.5 mL, 32 mmol) and trifluoromethyl-sulfonic acid (150 mL) in CH2Cl2 (10 mL) was added benzyl 2,2,2-trichloro-acetimidate (7.0 mL, 37 mmol). After stirring for several min, the mixture was poured into aqueous NaHCO3 and extracted with Et2O. The organic extract was washed with saturated aqueous NaCl, dried (MgSO4) and concentrated under reduced pressure. The residue was purified by flash chromatography, eluting with a solvent gradient of 9-17% EtOAc/Hex to afford the title compound: 1H NMR (CDCl3): δ 7.34 (m, 5H); 4.62 (s, 2H); 4.11 (s, 2H); 3.78 (s, 3H).

(b) Benzyl-N,N-dimethylglycolamide -

To dimethyl-amino(methyl)aluminum chloride [prepared from dimethylamine hydrochloride (3.4 g, 42 mmol) and trimethyl aluminum (21 mL of 2 M solution, 42 mmol)] in toluene (40 mL) was added methyl benzylglycolate (3.0 g, 17 mmol). After stirring at rt for 1.5 h, the mixture was poured into 3 N HCl and extracted with Et2O. The organic extract was washed with saturated aqueous NaCl, dried (MgSO4) and concentrated under reduced pressure. The residue was purified by flash chromatography, eluting with a solvent gradient of 9-50% EtOAc/Hex. The title compound was obtained as a colorless oil (1.2 g, 37%): 1H NMR (CDCl3): δ 7.4-7.1 (m, 5H); 4.61 (s, 2H); 4.18 (s, 2H); 2.98 (s, 3H); 2.95 (s, 3H).

(c) N,N-Dimethylglycolamide -

To a solution of benzyl-N,N-dimethylglycolamide (0.28 g, 1.5 mmol) in MeOH (5 mL) was added 10% palladium on activated carbon (0.15 g), and the resulting mixture was stirred under an atmosphere of H2. After 1 h, the mixture was filtered through a pad of Celite, and the filtrate was concentrated under reduced pressure to afford the title compound which was used without further purification: 1H NMR (CDCl3): δ 4.13 (s, 2H); 3.01 (s, 3H); 2.89 (s, 3H).

(d) N,N-Dimethyl-4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzoyl-oxyacetamide -

To a solution of 4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzoic acid (0.15 g, 0.42 mmol) [See Ex. 9 above] in DMF (10 mL) was added carbonyldiimidazole (0.34 g, 2.1 mmol). After stirring for 18 h at rt, N,N-dimethylglycolamide (0.43 g, 4.2 mmol) was added and stirring was continued for an additional 3h at rt. The reaction mixture was poured into H2O, extracted with EtOAc and evaporated. The residue was purified by flash chromatography eluting with
a solvent gradient of 2% MeOH/CHCl$_3$ to afford the title compound: CIMS (NH$_3$, m/z): 445 (M$^+$+H).

**Example 28** - **2-(4-Aminophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole** - A mixture containing 4-(4-fluorophenyl)-2-(4-nitrophenyl)-5-(4-pyridyl)-1H-imidazole (2.0 g, 5.6 mmol) [See Ex. 26 above] and 10% palladium on activated carbon (0.4 g) was stirred under an atmosphere of H$_2$ for 4 h, then was filtered through a pad of Celite. The filtrate was concentrated under reduced pressure, and the residue was purified by flash chromatography, eluting with a solvent gradient of 1-10% MeOH/CHCl$_3$. The title compound was obtained as a light orange solid (0.50 g, 27%): $^1$H NMR (DMSO-d$_6$): $\delta$ 8.40 (d, 2H); 7.73 (d, 2H); 7.57 (m, 2H); 7.35 (m, 4H); 6.62 (t, 2H).

**Example 29** - **4-(4-Fluorophenyl)-2-(4-methanesulfonamidophenyl)-5-(4-pyridyl)-1H-imidazole** - To a suspension of 2-(4-aminophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole (80 mg, 0.24 mmol) [See Ex. 28 above] and triethylamine (0.12 mL, 0.86 mmol) in CH$_2$Cl$_2$ (5 mL) was added methanesulfonyl chloride (55 mL, 0.72 mmol). After stirring at rt for 1 h, the mixture was poured into aqueous NaHCO$_3$ and extracted with EtOAc. The organic extract was washed with saturated aqueous NaCl, dried (MgSO$_4$) and concentrated under reduced pressure. The residue was purified by flash chromatography, eluting with a solvent gradient of 2-10% MeOH/CHCl$_3$ to afford the title compound as a tan solid (35 mg, 36%): ESMS (m/z): 409 (M$^+$+H).

**Example 30** - **4-(4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl)phenyl-sulfonamide** - **(a) Ethyl 4-sulfonamidobenzoate** - A solution of 4-carboxybenzenesulfonamide (5.0 g, 0.025 mol) in 20% ethanolic HCl (20 mL) was heated at reflux for 18 h, then allowed to cool and was concentrated under reduced pressure to afford the title compound: $^1$H NMR (CDCl$_3$): $\delta$ 8.20 (apparent d, 2H); 8.00 (apparent d, 2H); 4.88 (br s, 2H); 4.43 (q, 2H); 1.43 (t, 3H).

**(b) N-Methoxy-N-methyl-4-sulfonamidobenzamide** - To a solution of methoxymethylamino(methyl)aluminum chloride [prepared from methoxymethylamine hydrochloride (4.8 g, 50 mmol) and trimethyl aluminum (25 mL of 2 M solution, 50 mmol)] in toluene (50 mL) at 0 °C was added ethyl 4-sulfonamidobenzoate (3.8 g, 17 mmol). The mixture was allowed to warm to rt and stir for 3 h, then was poured into a slurry of silica gel (50 g) in CHCl$_3$ (200 mL). The mixture was filtered, and the filtrate was concentrated under reduced pressure. The residue was poured into H$_2$O,
and the solid which formed was collected by filtration, washed with H₂O
and dried in vacuo to afford the title compound (1.7 g, 42%): ¹H NMR
(CDCl₃/Methanol-d₄): δ 7.86 (d, 2H); 7.66 (d, 2H); 3.43 (s, 3H); 3.29 (s, 3H).

(c) 4-Formylbenzenesulfonamide - To a solution of N-
methoxy-N-methyl-4-sulfonamidobenzamide (1.0 g, 4.1 mmol) in THF (25
mL) at -78 °C was added LiAlH₄ (6.1 mL of 1 M solution in THF, 6.1 mmol).
After stirring at -78 °C for 30 min, the mixture was poured into a slurry of
silica gel (50 g) in CHCl₃ (200 mL). The mixture was filtered and the
filtrate was concentrated under reduced pressure. The residue was purified
by flash chromatography, eluting with a solvent gradient of 2-10%
MeOH/CHCl₃. The title compound was obtained as a white solid (0.12 g,
16%): ¹H NMR (CDCl₃/Methanol-d₄): δ 10.3 (s, 1H); 8.02 (d, 2H); 7.95 (d, 2H).

(d) 4-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-
yl]sulfonamide - The title compound was prepared using the same
procedure as described in Example 13, except using 4-formylbenzene-
sulfonamide: ESMS (m/z): 395 (M⁺+H).

Example 31 - N'-Cyanomethyl-N-[4-(fluorophenyl)-5-(4-pyridyl)-1H-
imidazol-2-yl]benzylguanidine - To a suspension of 2-(4-aminomethylphenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole (0.10 g,
0.29 mmol) [See Ex. 11 above] in CH₃CN was added diphenyl
cyanocarbonimide (83 mg, 0.35 mmol). After stirring at rt for 18 h, the
solid which formed was collected by filtration and washed with CH₃CN.
The solid was dissolved in MeOH saturated with NH₃ and stirred for 72 h.
The mixture was concentrated under reduced pressure, and the residue was
purified by flash chromatography, eluting with a solvent gradient of 4-10%
MeOH/CHCl₃. The title compound was isolated as a pale yellow solid (22
mg, 18%): mp 280-281 °C.

Example 32 - 2-[4-(Methanesulfonamido)methylphenyl]-4-(4-
fluorophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was
prepared using the same procedure as described in Example 29, except
using 2-(4-aminomethylphenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-
imidazole [See Ex. 11 above]: ESMS (m/z): 423 (M⁺+H).

Example 33 - 4-(4-Fluorophenyl)-2-(4-methoxyphenyl)-5-(4-pyridyl)-
1H-imidazole - (a) 1-Cyano-1-(4-pyridyl)methyl 4-methoxybenzoate -
The title compound was prepared using the same procedure Lantos, I. et al.
(J. Med. Chem. 1984, 27, 72-75) employed to prepare 1-cyano-1-(4-pyridyl)-
methyl benzoate, except using 4-methoxybenzoyl chloride: ¹H NMR
(CDCl₃): δ 8.81 (d, 2H); 8.10 (d, 2H); 7.57 (d, 2H); 7.01 (d, 2H); 6.74 (s, 1H); 3.93 (s, 3H).

(b) 1-(4-Fluorophenyl)-2-(4-pyridyl)-2-oxoethyl 4-methoxybenzoate and 2-(4-fluorophenyl)-1-(4-pyridyl)-2-oxoethyl 4-methoxybenzoate - The title compounds were prepared using the same procedure Lantos et al. (J. Med. Chem. 1984, 27, 72-75) used to prepare 1-(4-Fluorophenyl)-2-(4-pyridyl)-2-oxoethyl benzoate and 2-(4-fluorophenyl)-1-(4-pyridyl)-2-oxoethyl benzoate, except using 1-Cyano-1-(4-pyridyl)methyl 4-methoxybenzoate: ¹H NMR (faster eluting isomer, CDCl₃): δ 8.78 (d, 2H); 8.03 (br d, 2H); 7.73 (d, 2H); 7.53 (dd, 2H); 7.10 (apparent t, 2H); 6.93 (overlapping s and d, 3H); 3.85 (s, 3H); ¹H NMR (slower eluting isomer, CDCl₃): δ 8.66 (d, 2H); 8.04 (m, 4H); 7.46 (d, 2H); 7.15 (apparent t, 2H); 6.95 (overlapping s and d, 3H); 3.87 (s, 3H).

(c) 4-(4-Fluorophenyl)-2-(4-methoxyphenyl)-5-(4-pyridyl)-1H-imidazole - To a solution containing a mixture of 1-(4-fluorophenyl)-2-(4-pyridyl)-2-oxoethyl 4-methoxybenzoate and 2-(4-fluorophenyl)-1-(4-pyridyl)-2-oxoethyl 4-methoxy-benzoate (0.35 g, 0.96 mmol) in glacial acetic acid (7.5 mL) was added ammonium acetate (0.35 g, 4.5 mmol). The resulting mixture was heated at reflux for 18 h, then allowed to cool. The mixture was concentrated under reduced pressure, and the residue was purified by flash chromatography, eluting with a solvent gradient of 33-60% EtOAc/Hex. The material which was isolated was recrystallized from MeOH/CH₂Cl₂ to provide the title compound (65 mg, 20%) as an off-white solid: mp 264-265 °C.

Example 34 - 2-(4-Amino-3-iodophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - To a solution of 2-(4-aminophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole (50 mg, 0.15 mmol) [See Ex. 28 above] in glacial acetic acid (5 mL) was added a solution of ICl (24 mg, 0.15 mmol) in glacial acetic acid (1.5 mL). The resulting mixture was stirred at rt for 1 h, then poured into saturated aqueous Na₂S₂O₅. The pH was adjusted to neutral by the addition of 2.5 N NaOH and extracted with EtOAc. The organic extract was concentrated under reduced pressure, and the residue was purified by flash chromatography, eluting with a solvent gradient of 2-10% MeOH/CHCl₃. The material that was isolated was recrystallized from Et₂O/Hex to afford the title compound: ¹H NMR (CDCl₃): δ 8.42 (d, 2H); 8.18 (d, 1H); 7.68 (dd, 2H); 7.42 (m, 4H); 7.09 (t, 2H); 6.77 (d, 1H).
Example 35 - N-Benzyl-N-methyl-4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzamide - The title compound was prepared using the same procedure as described in Example 21, except using benzylmethylaminodimethyl aluminium and methyl 4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzoate [See Ex. 18 above]: mp 233-234 °C.

Example 36 - 2-[4-(N-Benzyl-N-methyl)aminomethylphenyl]-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 11, except using N-benzyl-N-methyl-4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzamide [See Ex. 35 above]: ESMS (m/z): 449 (M⁺+H).

Example 37a - 4-(4-Fluorophenyl)-N-1-hydroxy-2-(4-methylthiophenyl)-5-(4-quinolyl)imidazole - (a) 4-Fluoro-2-(4-quinolyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (b) except using 4-methylquinoline: ¹H NMR (CDCl₃): δ 8.87 (d, 1H); 8.10 (m, 3H); 7.88 (d, 1H); 7.74 (br t, 1H); 7.57 (br t, 1H); 7.20 (m, 3H); 4.73 (s, 2H).

(b) 4-Fluoro-2-hydroxyimino-2-(4-quinolyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (c) except using 4-fluoro-2-(4-quinolyl)acetophenone.

(c) 4-(4-Fluorophenyl)-N-1-hydroxy-2-(4-methylthiophenyl)-5-(4-quinolyl)imidazole - The title compound was prepared using the same procedure as described in Example 10, step (d) except using 4-fluoro-2-hydroxyimino-2-(4-quinolyl)acetophenone and 4-(methylthio)benzaldehyde: ¹H NMR (CDCl₃): δ 8.03 (m, 1H); 7.80 (br d, 2H); 7.52 (d, 1H); 7.40-7.10 (m, 5H); 6.81 (br m, 3H); 6.61 (apparent t, 2H), 2.48 (s, 3H).

Example 37b - 4-(4-Fluorophenyl)-2-(4-methylthiophenyl)-5-(4-quinolyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 1, except using 4-(4-fluorophenyl)-N-1-hydroxy-2-(4-methylthiophenyl)-5-(4-quinolyl)imidazole: ESMS (m/z): 412 (M⁺+H).

Example 38 - 4-(4-Fluorophenyl)-2-(4-methylsulfinylphenyl)-5-(4-quinolyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 20, except using 4-(4-fluorophenyl)-2-(4-methylthiophenyl)-5-(4-quinolyl)-1H-imidazole: ESMS (m/z): 428 (M⁺+H).

Example 39 - 4-(3-Chlorophenyl)-2-(4-methylsulfinylphenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 20, except using 4-(3-chlorophenyl)-2-(4-
methylthiophenyl)-5-(4-pyridyl)-1H-imidazole [See Ex. 40 below]: ESMS (m/z): 394 (M⁺+H).

Example 40a - 4-(3-Chlorophenyl)-N-1-hydroxy-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole - (a) 3-Chloro-N-methoxy-N-methylbenzamide - The title compound was prepared using the same procedure as described in Example 10 (a) except using 3-chlorobenzoyl chloride: 1H NMR (CDCl₃): δ 7.69 (br s, 1H); 7.58 (br d, 1H); 7.42 (br d, 1H); 7.31 (dd, 1H); 3.55 (s, 3H); 3.34 (s, 3H).

(b) 3-Chloro-2-(4-pyridyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (b) except using 4-picoline and 3-chloro-N-methoxy-N-methylbenzamide: 1H NMR (CDCl₃): δ 8.60 (d, 2H); 8.00 (br s, 1H); 7.89 (br d, 1H); 7.60 (br d, 1H); 7.45 (t, 1H); 7.21 (d, 2H); 4.27 (s, 2H).

(c) 3-Chloro-2-hydroxyimino-2-(4-pyridyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (c) except using 3-chloro-2-(4-pyridyl)acetophenone.

(d) 4-(3-Chlorophenyl)-N-1-hydroxy-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 10, step (d) except using 3-chloro-2-hydroxyimino-2-(4-pyridyl)acetophenone and 4-methylthiobenzaldehyde: 1H NMR (CDCl₃): δ 8.04 (d, 2H); 7.70 (d, 2H); 7.21-6.91 (m, 8H); 2.47 (s, 3H).

Example 40b 4-(3-Chlorophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 1, except using 4-(3-chlorophenyl)-N-1-hydroxy-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole: ESMS (m/z): 378 (M⁺+H).

Example 41 - 4-(4-Fluorophenyl)-2-(4-formamidomethylphenyl)-5-(4-pyridyl)-1H-imidazole - Formic acid (10 ml) was added to acetic anhydride (20 mL) and the mixture was heated to 50 °C for 15 min. 2-(4-Aminomethylphenyl)-4-(4-fluorophenyl)-5-4-pyridyl)imidazole (0.25 g, 0.73 mmol) [See Ex. 11] was added and the reaction mixture was heated to 50 °C for 2 h. The solvent was evaporated and the residue was purified by flash chromatography, eluting with a solvent gradient of 4-10% MeOH/CHCl₃. The title compound was isolated as a tan solid (0.15 g, 55%): ESMS (m/z): 373 (M⁺+H).

Example 42 - 4-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-y]benzohydroxamic acid - To a solution of O-benzyl-4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imazol-2-yl]benzohydroxamic acid (0.10 g, 0.22 mmol) [See Ex. 43 below] in ethanol (10 mL) was added 10% palladium on carbon. After stirring under an atmosphere of H₂ for 18 h, the reaction mixture was filtered through
celite and the solids were washed with ethanol. The combined filtrates were evaporated and the residue was recrystallized from 2-propanol to afford the title compound (0.040 g, 50%): ESMS (m/z): 375 (M+H).

Example 43 - O-Benzyl-4-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]-benzohydroxamic acid - To a stirred suspension of O-benzylhydroxylamine hydrochloride (1.2 g, 7.8 mmol) in toluene (20 mL) at 0 °C was added trimethylaluminum (2.0 M in toluene, 3.9 mL, 7.8 mmol). The reaction mixture was warmed to rt and stirring was continued at this temperature for 1 h. Ethyl 4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl] benzoate (1.0 g, 2.6 mmol) [See Ex. 14 above] was added and the reaction mixture was heated at reflux for 3 h. After cooling, the reaction was poured into 10% MeOH/CHCl₃ containing silica gel. The solids were removed by filtration and washed with 10% MeOH/CHCl₃. The combined washings were evaporated and the residue was purified by flash chromatography eluting with a solvent gradient of 1-10% MeOH/CHCl₃. Trituration with ether afforded the title compound as a white solid (0.25 g, 21%): ¹H NMR (CDCl₃/MeOH-d₄): δ 8.16 (d, 2H); 7.77 (d, 2H); 7.53 (d, 2H); 7.23 (m, 5H); 7.10 (m, 4H); 6.88 (t, 2H).

Example 44 - 4-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzamide oxime - To a mixture of 2-(4-cyanophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole (3.0 g, 8.7 mmol) [See Ex. 1 above] and K₂CO₃ (2.4 g, 17 mmol) in EtOH (120 mL) and H₂O (6 mL) was added hydroxylamine hydrochloride (1.2 g, 17 mmol). After heating at reflux for 24 h, the reaction mixture was poured into H₂O. The precipitate was collected, washed with H₂O and air-dried. The crude product was dissolved in acetone, silica gel was added and the solvent was evaporated. The impregnated silica gel was added to the top of a flash column and the column was eluted with a solvent gradient of 2-10% MeOH/CHCl₃ to afford the title compound as a white solid (3.0 g, 91%): ESMS (m/z): 374 (M+H).

Example 45 - N"-Methyl-N'-cyano-N-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzylguanidine - To a suspension of 2-(4-aminomethylphenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole (2.5 g, 7.3 mmol) [See Ex. 11 above] in CH₃CN (250 mL) was added diphenyl cyanocarbonimidate (8.8 g, 7.3 mmol). After stirring at rt for 18 h, the solid which formed was collected by filtration and washed with CH₃CN (2.1 g, 59%). Without further purification, this material was added to methanol (100 mL) saturated with methylamine. The flask was stoppered and the reaction was stirred for 18 h at rt. The solvent and excess methylamine were evaporated and the residue was triturated with ether to give a brown solid which was further purified by flash chromatography eluting with a solvent gradient of 4-
10% MeOH/CHCl₃ to afford the title compound as a tan solid (0.33 g, 78%): CIMS (NH₃, m/z): 426 (M⁺+H).

Example 46a - N-1-Hydroxy-4-(3-methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole - (a) 3-Methoxy-N-methoxy-N-methylbenzamide - The title compound was prepared using the same procedure as described in Example 10, step (a) except using m-anisoyl chloride: ¹H NMR (CDCl₃): δ 7.27 (m, 3H); 7.01 (m, 1H); 3.82 (s, 3H); 3.57 (s, 3H); 3.36 (s, 3H).

(b) 3-Methoxy-2-(4-pyridyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (b) except using 3-methoxy-N-methoxy-N-methylbenzamide: ESMS (m/z): 228.2 (M⁺+H).

(c) 2-Hydroxyimino-3-methoxy-2-(4-pyridyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (c) except using 3-methoxy-2-(4-pyridyl)acetophenone.

(d) N-1-Hydroxy-4-(3-methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 10, step (d) except using 2-hydroxyimino-3-methoxy-2-(4-pyridyl)acetophenone and 4-(methylthio)benzaldehyde: ESMS(m/z): 390 (M⁺+H).

Example 46b - 4-(3-Methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole - The title compound was prepared using the same procedure as described in Example 1, except using N-1-hydroxy-4-(3-methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole: mp 215.0-216.0 °C.

Example 47 - 4-(3-Methoxyphenyl)-2-(4-methylsulfanylphenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 38, except using 4-(3-methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole [See Example 46 above]: mp 167-168.5 °C.

Example 48 - Morpholino-4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzamide - The title compound was prepared using the same procedure as described in Example 21 except using dimethylamino-(morpholino)aluminum chloride and ethyl 4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]-benzoate [See Ex. 14 above]: ESMS (m/z): 429 (M⁺+H).

Example 49 - 4-(4-Fluorophenyl)-5-[4-(2-methylpyridyl)]-2-(4-methylthiophenyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 13, except using 4-(4-fluorophenyl)-1-hydroxy-5-[4-(2-methylpyridyl)]-2-(4-methylthiophenyl)-imidazole [See Ex. 66 below]: ESMS (m/z): 376.2 (M⁺+H).
Example 50 - 4-(4-Fluorophenyl)-5-{4-(2-methylpyridyl)]-2-(4-methylsulfonylphenyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 20 except using 4-(4-fluorophenyl)-5-[4-(2-methylpyridyl)]-2-(4-methylthiophenyl)-1H-imidazole [See Ex. 49 above]: ESMS (m/z): 392.2 (M^+H).

Example 51a - 4-(4-Fluorophenyl)-N-1-hydroxy-5-(4-pyrimidinyl)imidazole - (a) 4-Fluoro -2-(4-pyrimidinyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (b) except using 4-methylpyrimidine.

(b) 4-Fluoro-2-hydroxyimino-2-(4-pyrimidinyl)acetophenone - The title compound was prepared using the same procedure described in Example 10, step (c) except using 4-fluoro-2-(4-pyrimidinyl)acetophenone.

(c) 4-(4-Fluorophenyl)-N-1-hydroxy-5-(4-pyrimidinyl)imidazole - The title compound was prepared using the same procedure described in Example 10, step (d) except using 4-fluorophenyl-2-hydroxyimino-2-(4-pyrimidinyl)acetophenone.

Example 51b - 4-(4-Fluorophenyl)-2-(4-methylthiophenyl)-5-(4-pyrimidinyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 1, except using 4-(4-fluorophenyl)-N-1-hydroxy-5-(4-pyrimidinyl)imidazole: CIMS (NH₃, m/z): 363 (M^+H).

Example 52 - 4-(4-Fluorophenyl)-2-(4-methylsulfonylphenyl)-5-(4-pyrimidinyl)-1H-imidazole - The title compound was prepared using the same procedure described in Example 20, except using 4-(4-fluorophenyl)-2-(4-methylthio)phenyl)-5-(4-pyrimidinyl)-1H-imidazole: CIMS (NH₃, m/z): 379 (M^+H).

Example 53 - 4-(4-Fluorophenyl)-2-(4-methylsulfonylphenyl)-5-(4-pyrimidinyl)-1H-imidazole - To a solution of 4-(4-fluorophenyl)-2-(4-methylthiophenyl)-5-(4-pyrimidinyl)-1H-imidazole (0.10 g, 0.28 mmol) [See Ex. 51 above] was added 3-chloroperbenzoic acid (50%, 0.15 g, 0.42 mmol). After stirring at rt for 72 h, the solvent was evaporated and the residue was partitioned between EtOAc and 2.5 N NaOH. The organic phase was washed with brine, dried (MgSO₄) and evaporated. The residue was triturated with EtOAc to afford the title compound as a white solid (0.50 g, 46%). CIMS (NH₃, m/z): 395 (M^+H).

Example 54 - 4-(4-Fluorophenyl)-2-(4-Morpholinomethylphenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 11 except using morpholino-4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzamide: mp 242-243 °C.

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Example 55 - 4-(4-Fluorophenyl)-2-(4-hydroxymethyl)-5-(4-pyridyl)-1H-imidazole - To a suspension of ethyl 4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl] benzoate (1.0 g, 2.6 mmol) [See Ex. 14 above] in THF (25 mL) was added LiAlH₄ (1 M in THF, 7.8 mL, 7.8 mmol). After stirring at rt for 0.5 h, the reaction mixture was poured into 2.5 N NaOH and extracted three times with 2:1 EtOAc/CH₂Cl₂. The combined extracts were washed with brine, dried (MgSO₄) and evaporated to afford the title compound as a white solid (0.50 g, 54%).

Example 56 - 4-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzaldehyde - To a suspension of 4-(4-fluorophenyl)-2-(4-hydroxymethyl)-5-(4-pyridyl)-1H-imidazole (0.40 g, 1.2 mmol) [See Ex. 55 above] in CH₂Cl₂ (40 mL) was added pyridinium chlorochromate (0.30 g, 1.4 mmol) at rt. The reaction mixture was stirred at this temperature for 2 h, filtered through a pad of silica gel eluting with 2% MeOH/CHCl₃ and the filtrate evaporated. The residue was purified by flash chromatography eluting with 4% MeOH/CHCl₃ followed by recrystallization from CH₂Cl₂/MeOH to afford the title compound as a white solid (0.30 g, 7.3%).

Example 57 - 4-(2-Methoxyphenyl)-2-(4-methylsulfinylphenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 20 except using 4-(2-methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole [See Ex. 58 below]: CIMS (NH₃,m/z): 390 (M^+ + H).

Example 58a - N-1-Hydroxy-4-(2-methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole - (a) 2-Methoxy-N-methoxy-N-methylbenzamide - The title compound was prepared using the same procedure as described in Example 10, step (a) except using o-anisoyl chloride: ^1H NMR (CDCl₃): δ 7.36 (m, 3H); 6.98 (dd, 1H); 3.84 (s, 3H); 3.56 (br s, 3H); 3.32 (br s, 3H).

(b) 2-Methoxy-2-(4-pyridyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (b) except using 2-methoxy-N-methoxy-N-methylbenzamide.

(c) 2-Hydroxyimino-2-methoxy-2-(4-pyridyl)acetophenone - The title compound was prepared using the same procedure as described in Example 10, step (c) except using 2-methoxy-2-(4-pyridyl)acetophenone.

(d) N-1-Hydroxy-4-(2-methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole - The title compound was prepared using the same procedure as described in Example 10, step (d) except using 2-hydroxyimino-2-methoxy-2-(4-pyridyl)acetophenone and 4-(methylthio)benzaldehyde: ESMS (m/z): 390.0 (M^+ + H).
Example 58b - 4-(2-Methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 1, except using N-1-hydroxy-4-(2-methoxyphenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole: CIMS (NH₃, m/z): 374.2 (M⁺+H).

Example 59 - 3-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]phenyl-5-methyl-4,5-dihydro-1,2,4-oxadiazole - A solution of 4-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzamide oxime (0.50 g, 1.3 mmol) [See Ex. 44 above] and acetaldehyde (25 mL) in ethanol (100 mL) and H₂O (100 mL) was stirred at rt for seven days. The solvent was evaporated and the residue was purified by flash chromatography eluting with a solvent gradient of 2-4% CHCl₃/MeOH. Recrystallization from EtOAc afforded the title compound as a yellow solid (0.11 g, 21%): CIMS (NH₃, m/z): 400 (M⁺+H).

Example 60 - 3-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]phenyl-5-methyl-1,2,4-oxadiazole - To a solution of 4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzamide oxime (0.10 g, 0.27 mmol) [See Ex. 44 above] in pyridine (10 mL) was added acetic anhydride (1.0 mL) at rt. After stirring at this temperature for 18 h, the reaction mixture was poured into H₂O, and the precipitate collected, washed with H₂O and dried in vacuo.

Without further purification, the crude o-acylamidoxime was dissolved in acetic acid (5 mL) and heated at reflux for 3 h. The solvent was evaporated, H₂O was added and the mixture was neutralized with aqueous NaHCO₃. The precipitate was collected, washed with H₂O, air-dried and purified by flash chromatography eluting with 4% MeOH/CHCl₃. Trituration with ether afforded the title compound as a white solid (0.030 g, 28%): CIMS (NH₃, m/z): 398 (M⁺+H).

Example 61 - 4-(3-Aminophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole - A solution of 0.161 g (0.41 mmol) of 2-(4-methylthiophenyl)-4-(3-nitrophenyl)-5-(4-pyridyl)-1H-imidazole [See Ex. 62 below] in 3.4 mL of HOAc-H₂O (1:1) was treated with 1.81 mL (2.87 mmol) of 20% aqueous titanium (III) chloride in one single portion. The mixture was stirred at rt for 20 min, then made basic with 10% NaOH. The aqueous mixture was extracted with 95:5 CH₂Cl₂/MeOH. The organic extracts were washed with H₂O and saturated NaCl. Evaporation of solvent in vacuo afforded a yellow solid which was filtered through a plug of silica gel, eluting with 90:10 CHCl₃/MeOH. The title compound was isolated as a pale yellow solid (0.129 g, 78%): CIMS (NH₃, m/z): 359.1 (M⁺+H).

Example 62a - N-1-Hydroxy-2-(4-methylthiophenyl)-4-(3-nitrophenyl)-5-(4-pyridyl)imidazole - (a) 1-(3-Nitrophenyl)-2-(4-pyridyl)ethanol - The
title compound was prepared using the same procedure as described in
Example 10, step (b) except using 3-nitrobenzaldehyde: 1H NMR (CDCl₃): δ
8.41 (d, 2H); 8.23 (s, 1H); 8.15 (d, 1H); 7.67 (d, 1H); 7.54 (t, 1H); 7.19 (d, 2H);
5.05 (t, 1H); 4.41 (s, 2H).

(b) 1-(3-Nitrophenyl)-2-(4-pyridyl)acetophenone - To a solution of
1.0 mL (14.3 mmol) of DMSO in 55 mL of dry CH₂Cl₂ was added 1.82 mL (12.9
mmol) of trifluoroacetic anhydride at -78 °C. The mixture was stirred for 30
min, then a solution of 1-(3-nitrophenyl)-2-(4-pyridyl)ethanol (1.09 g, 4.46
mmol) in DMSO/CH₂Cl₂ (3/11 mL) was added dropwise. The mixture was
stirred at -78 °C for 2 h, then 4.1 mL (29.4 mmol) of triethylamine was added
dropwise. The ice bath was removed and the mixture was warmed to room
temperature. The mixture was poured into saturated NH₄Cl and extracted
with CH₂Cl₂. The organic extracts were washed with saturated NH₄Cl and
saturated NaCl, then dried over MgSO₄. Removal of the solvent in vacuo
afforded a red oil which was purified by flash chromatography, eluting with a
gradient of 0-3% MeOH/CHCl₃. The title compound was isolated as an orange
oil (0.65 g, 60%): 1H NMR (CDCl₃): δ 8.83 (s, 1H); 8.60 (d, 2H); 8.46 (d, 1H);
8.32 (d, 1H); 7.72 (t, 1H); 7.23 (d, 2H); 4.38 (s, 2H).

c) 2-Hydroxyimino-1-(3-nitrophenyl)-2-(4-pyridyl)acetophenone
The title compound was prepared using the same procedure as described in
Example 10, step (c) except using 1-(3-nitrophenyl)-2-(4-pyridyl)acetophenone.

d) N-1-Hydroxy-2-(4-methylthiophenyl)-4-(3-nitrophenyl)-5-(4-
pyridyl)imidazole - The title compound was prepared using the same
procedure as described in Example 10, step (d) except using 2-hydroxyimino-1-
(3-nitrophenyl)-2-(4-pyridyl)acetophenone and 4-(methylthio)benzaldehyde: 1H
NMR (CDCl₃/MeOH-d₄): δ 8.55 (d, 2H); 8.43 (m, 1H); 8.15 (dd, 1H); 8.06 (d,
2H); 7.78 (d, 1H); 7.51 (m, 1H); 7.45 (d, 2H); 7.32 (m, 2H); 2.57 (s, 3H).

Example 62b - 2-(4-Methylthiophenyl)-4-(3-nitrophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 1, except using N-1-hydroxy-2-(4-methylthiophenyl)-4-(3-
nitrophenyl)-5-(4-pyridyl)imidazole: CIMS (NH₃, m/z): 389.1 (M⁺+H)

Example 63 - 4-(3-Methanesulfonylamidophenyl)-2-(4-methylthiophenyl)-
5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same
procedure as described in Example 29, except using 4-(3-aminophenyl)-2-
(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole [See Ex. 61 above]: ESMS
(m/z): 437.0 (M⁺+H).

Example 64 - 3-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-
yl]phenyl-1,2,4-oxadiazol-5(4H)-one - To a mixture of 4-[4-(4-fluorophenyl)-
5-(4-pyridyl)-1H-imidazol-2-yl]benzamide oxime (0.25 g, 0.67 mmol) [See Ex. 44
above) in CH2Cl2 (5.0 mL) and Et3N (0.19 mL, 1.3 mmol) was added ethyl chloroformate (0.076 mL, 0.80 mmol) at rt. After 0.5 h at this temperature, the reaction mixture was poured into H2O, extracted four times with CH2Cl2 and once with 10% MeOH/CH2Cl2. The organic extracts were combined and evaporated. The residue was purified by flash chromatography eluting with a solvent gradient of 2-4% MeOH/CHCl3. Trituration with ether afforded a white solid (0.22 g, 73%). A portion of this compound (0.10 g, 0.22 mmol) was dissolved in HOAc (2.5 mL) and heated to reflux for 18 h. The reaction mixture was poured into H2O, neutralized with concentrated NH4OH, extracted with EtOAc and evaporated. The residue was triturated sparingly with cold EtOAc to afford the title compound as a yellow solid (0.020 g, 23%): CIMS (NH3, m/z): 400 (M+H).

Example 65 - 4-(3-Acetamidophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 6, except using 4-(3-aminophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole [See Ex. 61 above]: ESMS (m/z): 401 (M+H).

Example 66 - 4-(4-Fluorophenyl)-1-N-hydroxy-5-[4-(2-methylpyridyl)]-2-(4-methylthiophenyl)imidazole - (a) 2-Methyl isonicotinic acid - The title compound was prepared using the same procedure as described in Liebig's Ann. Chem., 1958, 613, 153: ESMS (m/z): 138.0 (M+H).

(b) Methyl 2-methylisonicotinate - To an ice-cooled suspension of 1.32 g (9.62 mmol) of 2-methylisonicotinic acid in 20 mL of MeOH was added 1.47 mL (20.2 mmol) of thionyl chloride. The ice-bath was removed and the reaction was stirred at rt. After 22 h, the MeOH was evaporated and the residue was taken up in H2O. The aqueous mixture was neutralized with saturated NaHCO3, then extracted with Et2O. The organic extracts were washed with saturated NaCl, dried over MgSO4, then filtered through a bed of celite. Evaporation of solvent in vacuo afforded the title compound as a yellow liquid (0.89 g, 61%): 1H NMR (CDCl3): d 8.66 (d, 1H); 7.72 (s, 1H); 7.64 (d, 1H); 3.98 (s, 3H); 2.64 (s, 3H).

(c) Methyl 4-fluorophenylacetate - The title compound was prepared using the same procedure as described in Example 66, step (b) except using 4-fluorophenylacetic acid: 1H NMR (CDCl3): d 7.25 (dd, 2H); 7.02 (t, 2H); 3.71 (s, 3H); 3.61 (s, 2H).

(d) 2-(4-Fluorophenyl)-1-[2-methyl-(4-pyridyl)]ethanone - To a freshly prepared solution of NaOMe (3.0 M in MeOH) was added a solution of methyl 2-methylisonicotinate (6.81 g, 45.1 mmol) in MeOH (10 mL). This was
followed by the dropwise addition of a solution of methyl 4-fluorophenylacetate (8.34 g, 49.6 mmol) in MeOH (10 mL). The MeOH was distilled off while heating the reaction mixture at 95 °C. After 17.5 h, the solid residue was cooled. Concentrated HCl (15 mL) was added, and the mixture was heated at reflux. After 4 h, the mixture was cooled then diluted with H2O. The aqueous mixture was washed with Et2O, adjusted to pH 5 with 1N NaOH, then adjusted to pH 8 with saturated NaHCO3. The alkaline aqueous was extracted with EtOAc. The EtOAc extracts were washed with saturated NaCl, then dried over Na2SO4. Evaporation of solvent in vacuo afforded a red oil which was purified by column chromatography, eluting with a gradient of 0-3% MeOH/CHCl3. The title compound was isolated as a red oil (1.5 g, 15%).

(e) 2-(4-Fluorophenyl)-2-hydroxyimino-1-[2-methyl-(4-pyridyl)]ethanone - The title compound was prepared using the same procedure as described in Example 10, step (c) except using 2-(4-fluorophenyl)-1-[2-methyl-(4-pyridyl)]ethanone: ESMS (m/z): 259 (M+H).

(f) 4-(4-Fluorophenyl)-1-hydroxy-5-[4-(2-methylpyridyl)]-2-(4-methylthiophenyl)imidazole - The title compound was prepared using the same procedure as described in Example 10, step (d) except using 2-(4-fluorophenyl)-2-hydroxyimino-1-[2-methyl-(4-pyridyl)]ethanone and 4-(methylthio)benzaldehyde: ESMS (m/z): 392 (M+H).

Example 67 - 3-[4-(4-Fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]-phenyl-5,5-dimethyl-4,5-dihydro-1,2,4-oxadiazole - To a solution of 4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzamide oxime (0.25 g, 0.67 mmol) [See Ex. 44 above] in acetone (10 mL) was added pyridinium trifluoroacetate (0.39 g, 2.0 mmol). After heating at reflux for 18 h, the reaction mixture was poured into saturated aqueous NaHCO3, extracted with EtOAc and the organic phase was evaporated. The residue was purified by flash chromatography eluting with a solvent gradient of 2-10% MeOH/CHCl3 to afford the title compound as a white solid (0.12 g, 43 %): CIMS (NH3, m/z): 414 (M+H).

Example 68 - N-Hydroxy-N-1-[4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]phenyl]ethyl urea

(a) a-Methyl-4-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]benzyl alcohol To a mixture of 2-(4-cyanophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole (1.0 g, 2.9 mmol) [See Ex. 1 above] was added MeMgBr (3 M in Et2O, 4.0 mL, 12 mmol) at rt. The reaction mixture was heated at reflux for 1 h, poured into saturated aqueous NH4Cl, extracted with THF and the organic phase was evaporated. The residue was dissolved in MeOH (20 mL) and NaBH4 (1.0 g, 26 mmol) was added. After 0.5
h at rt, the solvent was evaporated and the residue was purified by flash chromatography eluting with a solvent gradient of 1-10% MeOH/CHCl₃ to afford the title compound as a white solid (0.26 g, 25%): ¹H NMR (CDCl₃/MeOH-d₄): δ 8.37 (d, 2H); 7.79 (d, 2H); 7.4-7.2 (m, 6H); 6.99 (t, 3H); 4.76 (q, 1H); 1.35 (d, 3H).

(b) N-Hydroxy-N-[1-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]phenyl]ethylurea - To a mixture of a-Methyl-4-(4-fluorophenyl)-5-(4-pyridyl)imidazol-2-yl benzyl alcohol (0.25 g, 0.70 mmol), P(Ph)₃ (0.46 g, 1.75 mmol) and N,O-bis(benzzyloxy carbonyl)hydroxylamine (0.48 g, 1.75 mmol) in THF (15 mL) was added DEAD (0.28 mL, 1.75 mmol) at rt. The reaction mixture was stirred at this temperature for 3 h and the solvent evaporated. The residue was partially purified by flash chromatography eluting with 1% MeOH/CHCl₃. Methanol (25 mL) was added to this material and the mixture was cooled to -78 °C. Ammonia was bubbled in at this temperature for 15 min. The reaction mixture was warmed slowly to rt, stoppered and stirred at rt for 2 days. The solvent was evaporated and the residue was purified by flash chromatography eluting with a solvent gradient of 1-10% MeOH/CHCl₃. The title compound was obtained as an off-white solid (0.43 g, 14%): FABMS (m/z): 418 (M⁺+H).

Example 69 - N-Hydroxy-N-[1-[4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazol-2-yl]phenyl]methyl urea - The title compound was obtained using the same procedure described in Example 68, except using 4-(4-fluorophenyl)-2-(4-hydroxymethyl)-5-(4-pyridyl)-1H-imidazole: FABMS (m/z): 418 (M⁺+H).

Example 70 - 4-(3-Methylthiophenyl)-2-(4-morpholinomethylphenyl)-5-(4-pyridyl)-1H-imidazole -

(a) 3-Methylthiobenzaldehyde - The title compound was prepared using the same procedure as described by Campbell, J. R. in J. Org. Chem., 1962, 27, 2207: ¹H NMR (CDCl₃): δ 9.95 (s, 1H); 7.72 (s, 1H); 7.61 (d, 1H); 7.45 (m, 2H); 2.53 (s, 3H).

(b) 1-(3-Methylthiophenyl)-2-(4-pyridyl)ethanol - The title compound was prepared using the same procedure as described in Example 10, step (b) except using 3-(methylthio)benzaldehyde: ¹H NMR (CDCl₃): δ 8.33 (d, 2H); 7.0-7.5 (m, 6H), 4.87 (m, 1H); 2.96 (m, 2H); 2.45 (s, 3H).

(c) 1-(3-Methylthiophenyl)-2-(4-pyridyl)ethanedione - To a solution of 1-(3-methylthiophenyl)-2-(4-pyridyl)ethanol (2.5 g, 10.2 mmol) in CH₂Cl₂ (150 mL) was added a mixture of celite (4.4 g) and pyridinium dichromate (4.4 g, 20.4 mmol). After stirring for 12 h, the mixture was filtered through celite. The solvent was removed in vacuo, and the residue was purified by flash chromatography, eluting with a solvent gradient of 40-50% EtOAc/Hex to
provide the title compound (144 mg, 5.5%): ¹H NMR (CDCl₃): δ 8.88 (br d, 2H); 7.85 (s, 1H); 7.78 (d, 2H); 7.67 (d, 1H); 7.56 (d, 1H); 7.44 (t, 1H); 2.55 (s, 3H).

(d) 4-Morpholinomethylbenzaldehyde diethyl acetal - The title compound was prepared using the same procedure as described by Borch, R. F., Bernstein, M. D., and Durst, H. D. in J. Am. Chem. Soc., 1971, 93, 2897 except using the diethyl acetal: ¹H NMR (CDCl₃): δ 7.41 (d, 2H); 7.32 (d, 2H); 5.48 (s, 1H); 3.3-3.8 (m, 10H); 2.43 (br s, 4H); 1.25 (t, 6H).

(e) 4-(3-Methylthiophenyl)2-(4-morpholinomethylphenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 13 except using 1-(3-methylthiophenyl)-2-(4-pyridyl)-ethanedione and 4-morpholinomethylbenzaldehyde diethyl acetal: ¹H NMR (CDCl₃): δ 8.47 (d, 2H); 8.02 (d, 2H); 7.3-7.9 (m, 8H); 3.72 (t, 4H); 3.54 (s, 2H); 2.44 (br s, 4H); 2.38 (s, 3H).

Example 71 - 4-(3-Methylenesulfinylphenyl)-2-(4-morpholinomethylphenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described for Example 20, except using 4-(3-methylthiophenyl)2-(4-morpholinomethylphenyl)-5-(4-pyridyl)-1H-imidazole [See Ex. 70 above]: ¹H NMR (CDCl₃): δ 8.38 (d, 2H); 7.92 (d, 2H); 7.1-7.6 (m, 8H); 3.76 (t, 4H); 3.59, (s, 2H); 2.73 (br s, 4H).

Example 72 - 4-(3-Methanesulfonamidophenyl)-2-(4-methylenesulfinylphenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 20, except using 4-(3-methanesulfonamido-phenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole [See Ex. 63 above]: CIMS (NH₃,m/z): 453.3 (M⁺+H).

Example 73 - 2-(4-Ethylthiophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 13, except using 4-ethylthiobenzaldehyde: mp 203-205 °C.

Example 74 - 2-(4-Ethylsulfinylphenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 20, except using 2-(4-ethylthiophenyl)-4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole [See Ex. 73 above]: mp 240 °C.

Example 75 - 4-(4-Fluorophenyl)-2-[(4-(4-methyl-1-piperziny1)-sulfonyl-phenyl]-5-(4-pyridyl)-1H-imidazole

(a) Ethyl [4-(4-methyl piperaziny1) sulfonamido] benzoate - A mixture of 4-chlorosulfonyl benzoic acid (5.0 g, 22.67 mmol), N-methyl
piperazine (25 mL) and MeOH (5 mL) was stirred for 18 h and ether (200 mL) was added to the mixture. The crystalline solid precipitate was filtered and washed with ether (200 mL). The solid was suspended in 20% ethanolic HCl and the mixture was heated at reflux until a homogeneous solution was attained (about 2 h). The solution was cooled to rt, concentrated, and the residue was partitioned between sat. NaHCO₃ and EtOAc. The organic extract was dried and concentrated to yield the title compound (5.8 g 80%).

(b) 4-(4-Methyl piperazinyl) sulfonamido benzyl alcohol - The title compound was prepared using the same procedure as described in Example 78, step (b) except using ethyl [4-(4-methyl piperazinyl)sulfonamido] benzoate.

(c) 4-(4-Methyl piperazinyl) sulfonamido benzaldehyde - To a solution of oxalyl chloride (1.06 mL, 12.1 mmol) in CH₂Cl₂ (20 mL) was added DMSO (1.8 mL, 25.4 mmol.) at -60 °C and the mixture was stirred for 25 min. A solution of 4-(4-methyl piperazinyl) sulfonamido benzyl alcohol (3.0 g, 10.5 mmol) in CH₂Cl₂ (25 mL) and DMSO (5 mL) was added and the mixture was stirred for 1.5 h at -60 °C. Triethylamine (7.4 mL) was added and the mixture was partitioned between brine and EtOAc. The organic extract was concentrated, then purified by flash chromatography to yield the title compound (1.0 g, 33%).

(d) 4-(4-Fluorophenyl)-2-[4-(4-methyl piperazinyl)sulfonamido phenyl]-5-(4-pyridyl)-1H-imidazole - The title compound was prepared using the same procedure as described in Example 13, except using 4-(4-methyl piperazinyl) sulfonamido benzaldehyde: mp 74-76 °C.

Example 76 - 4-(4-Fluorophenyl)-2-[4-(N-methylmethanesulfonamido)methylphenyl]-5-(4-pyridyl)-1H-imidazole

(a) Methyl 4-[(methanesulfonamido)methyl]benzoate - To a suspension of 4-(aminomethyl)benzoic acid (10 g, 66 mmol) in MeOH (100 mL) at 0 °C was added SOCl₂ (5.3 mL, 73 mmol) dropwise. The ice bath was removed and the reaction stirred at rt overnight. After heating the reaction at reflux for 4 h, the solvent was evaporated. The residue was suspended in CH₂Cl₂ (100 mL) at 0 °C and triethylamine (25 mL) was added, followed by the dropwise addition of methanesulfonyl chloride (7.75 mL, 100 mmol). The reaction was stirred at rt for 1 h, poured into ice H₂O, extracted with CH₂Cl₂, dried over anhydrous Na₂SO₄, filtered and evaporated. The crude product was flash chromatographed on silica gel eluting with 1% MeOH/CHCl₃. The title compound was isolated as a white solid (11.8 g, 74 %): ¹H NMR (CDCl₃): δ 8.03 (d, 2H); 7.42 (d, 2H); 4.9 (br t, 1H); 4.38 (d, 2H); 3.92 (s, 3H); 2.89 (s, 3H).
(b) Methyl 4-[(N-Methylmethanesulfonamido)methyl]benzoate
- To a mixture of methyl 4-[(methanesulfonamido)methyl]benzoate (5 g, 20.6 mmol) in MeOH (100 mL) at rt was added K₂CO₃ (2.9 g, 21 mmol). Methyl iodide (7 mL, 16 g, 112 mmol) was added and the mixture stirred overnight. The reaction was filtered and the solid washed with CHCl₃/MeOH. The combined filtrates were evaporated and the residue was purified by flash chromatography eluting with 0-5% MeOH/CHCl₃. The title compound was isolated as a white solid (4.9 g, 94%): ¹H (CDCl₃): δ 8.08 (d, 2H); 7.48 (d, 2H); 4.41 (s, 1H); 3.97 (s, 3H); 3.91 (s, 3H); 2.83 (s, 3H).

(c) 4-[(N-Methylmethanesulfonamido)methyl]benzyl alcohol
- The title compound was prepared using the same procedure as described in Example 78, step (b) except using methyl 4-[(N-methylmethanesulfonamido)methyl]benzoate: ¹H NMR (CDCl₃): δ 7.34 (m, 4H); 4.68 (s, 2H); 4.29 (s, 2H); 2.83 (s, 3H); 2.74 (s, 3H).

(d) 4-[(N-Methylmethanesulfonamido)methyl]benzaldehyde
- The title compound was prepared using the same procedure as described in Example 78, step (c) except using 4-[(N-methylmethanesulfonamido)methyl]benzyl alcohol: ¹H NMR (CDCl₃): δ 10.02 (s, 1H); 7.9 (d, 2H); 7.54 (d, 2H); 4.4 (s, 2H); 2.9 (s, 3H); 2.81 (s, 3H).

(e) 4-(4-Fluorophenyl)-2-[4-(N-methylmethanesulfonamido)methylphenyl]-5-(4-pyridyl)-1H-imidazole
- The title compound was prepared using the same procedure as described in Example 13, except using 4-[(N-methylmethanesulfonamido)methyl]benzaldehyde: mp 222-224 °C.

Example 77 - Diethyl [1-methyl-4-phenyl-5-(4-pyridyl)-imidazol-2-yl]methoxy]methylphosphonate

(a) N-Methyl-N-[4-picoly]formamide
- To a solution of 4-picoly chloride-HCl (15 g, 91.4 mmol) and N-methylformamide (53.4 mL, 914 mmol) in 300 mL of THF was added 80% NaH in mineral oil (5.48 g, 183 mmol). After stirring at rt for 18 h the mixture was quenched with ice water and partitioned between CH₂Cl₂ and H₂O. The organic extract was washed with aqueous NaCl and dried over MgSO₄. The solvent was removed in vacuo, and the residue was purified by flash chromatography, eluting with 50:1 CH₂Cl₂/MeOH. The title compound was obtained as a pale yellow oil (10.5 g, 76%): ESMS (m/z): 151 (M⁺+H).

(b) 1-Methyl-4-phenyl-5-[4-pyridyl]imidazole
- To a solution of diiso-propyl-amine (11.2 mL, 79.9 mmol) in 150 mL of THF at -78 °C was added n-butyllithium (31.9 mL of 2.5 M solution, 79.9 mmol). To the resulting mixture was added a solution of N-methyl-N-[4-picoly]formamide (10 g, 66.5 mmol) in THF. The resulting orange-brown solution was stirred
at -78 °C for 20 min, at which time benzonitrile (13.6 mL, 133 mmol) was added. The resulting dark brown mixture was allowed to warm to rt and stirred for 1 h, heated to reflux for 4 h, and then cooled to rt and partitioned between CH₂Cl₂ and H₂O. The organic extract was washed with aqueous NaCl and dried (MgSO₄). The solvent was removed in vacuo, and the residue was purified by flash chromatography, eluting with 50:1 CH₂Cl₂/MeOH. The title compound was obtained as a light tan solid (5.83 g, 37%): mp 158-159 °C.

(c) 2-Formyl-1-methyl-4-phenyl-5-[4-pyridyl]imidazole - To a solution of 1-methyl-4-phenyl-5-[4-pyridyl]imidazole (0.275 g, 1.17 mmol) in THF at -78 °C was added n-butyllithium (0.56 mL of 2.5 M solution, 1.40 mmol). The resulting red-orange solution was allowed to stir at -78 °C for 0.5 h when DMF (0.18 mL, 2.34 mmol) was added. The mixture was allowed to warm to rt and stir for 4 h, then quenched with ice water and partitioned between CH₂Cl₂ and H₂O. The organic extract was washed with aqueous NaCl and dried (MgSO₄). The solvent was removed in vacuo and the residue was purified by flash chromatography eluting with 50:1 CH₂Cl₂/MeOH. The title compound was obtained as a white solid (0.187 g, 61%): mp 167-168 °C.

(d) 2-Hydroxymethyl-1-methyl-4-phenyl-5-(4-pyridyl)imidazole - To a solution of 2-formyl-1-methyl-4-phenyl-5-[4-pyridyl] imidazole (0.830 g, 3.15 mmol) in MeOH at 0 °C was added NaBH₄ (0.143 g, 3.78 mmol). The mixture was stirred at rt for 0.5 h when the solvent was evaporated in vacuo and the residue was partitioned between CH₂Cl₂ and H₂O. The organic extract was washed with aqueous NaCl and dried (MgSO₄). The solvent was removed in vacuo and the residue was purified by flash chromatography eluting with 25:1 CH₂Cl₂/MeOH. The title compound was obtained as a white solid (0.608 g, 73%): mp 236-238 °C.

(e) Diethyl [1-methyl-4-phenyl-5-(4-pyridyl)-imidazol-2-yl]methoxy]-methyl-phosphonate - To a suspension of 80% NaH in mineral oil (0.013 g, 0.452 mmol) in DMF at 0 °C was added 1-methyl-2-hydroxymethyl-4-phenyl-5-[4-pyridyl] imidazole (0.100 g, 0.377 mmol) in DMF. The resulting bright yellow solution was stirred at 0 °C for 0.5 h when diethyl chloromethylphosphonate (0.070 mL, 0.452 mmol) dissolved in 0.079 mL of HMPA was added. The resulting mixture was stirred at 0 °C for 15 min and then warmed to rt. After 5 h, the solution was partitioned between CH₂Cl₂ and H₂O. The organic extract was washed with aqueous NaCl and dried (MgSO₄). The solvent was removed in vacuo, and the residue was purified by flash chromatography, eluting with 50:1 CH₂Cl₂/MeOH. The title compound was obtained as a light amber oil (0.088 g, 56%): ESMS (m/z): 416 (M⁺+H).
Example 78 - 4-(4-Fluorophenyl)-2-[4-(methanesulfonamido)methylphenyl]-5-(4-pyridyl)-1H-imidazole - (a) Methyl 4-[(methanesulfonamido)methyl]benzoate - The title compound was prepared using the same procedure as described in Example 76, step (a).

(b) 4-[(Methanesulfonamido)methyl]benzyl alcohol - To a mixture of methyl 4-[(methanesulfonamido)methyl]benzoate (3.6 g, 15 mmol) in THF (150 mL) was added LiAlH₄ (1 M in THF, 30 mL, 30 mmol). The reaction mixture was stirred at rt for 1 h and poured into 10% MeOH/CHCl₃ containing silica gel. The solids were removed by filtration, washed with 10% MeOH/CHCl₃ and the combined washings were evaporated to yield the title compound as a white solid (2.6 g, 80%).

c) 4-[(Methanesulfonamido)methyl]benzaldehyde - To a solution of 4-[(methanesulfonamido)methyl]benzyl alcohol (1.0 g, 4.6 mmol) in CH₂Cl₂ (25 mL) was added pyridinium chlorochromate (1.5 g, 7.0 mmol). The reaction mixture was stirred for 1 h at rt and poured through a pad of silica gel eluting with 2% MeOH/CHCl₃. The title compound was isolated as a tan solid (1.0 g, 100%): ¹H NMR (CDCl₃): δ 10.03 (s, 1H); 7.88 (d, 2H); 7.57 (d, 2H); 4.79 (br s, 1H); 4.43 (d, 2H); 2.93 (s, 3H).

d) 4-(4-Fluorophenyl)-2-[4-(methanesulfonamido)methylphenyl]-5-(4-pyridyl)-1H-imidazole - The title compound [also prepared in Example 32] was prepared using the same procedure as described in Example 13, except using 4-[(methanesulfonamido)methyl]benzaldehyde.

Example 79 - 4-(4-Fluorophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole

(a) 1-(t-Butyldimethylsilyloxy)-2-(4-fluorophenyl)-1-(4-pyridyl)ethanone - To a -20 °C solution of diisopropylamine (64.4 mL, 0.46 mol) and THF (120 mL) was added 207.8 mL (0.52 mol, 2.5 M solution in hexanes) of n-butyllithium dropwise over 15 min. The temperature was lowered to -15 °C and the mixture was stirred for 0.5 hr. The solution was cooled to -20 °C and 98.14 g (0.44 mol) of 4-(t-butyldimethylsilyloxy)methyl pyridine was added dropwise over 20 min. After stirring at -20°C for 45 min, a solution of 4-fluoro-N-methoxy-N-methylbenzamide (84.5 g, 0.46 mol) [See Ex. 10, step (a)] in THF (90 mL) was added dropwise over 0.5 hr. Once the addition was complete, the ice bath was removed and the reaction mixture was warmed to 0 °C for 1 hr, then stirred at rt for 1.5 hr. The mixture was poured into a solution of NH₄Cl (98 g) and H₂O (500 mL), then extracted with EtOAc (3 x 250 mL). The EtOAc extracts were washed with H₂O and saturated NaCl, then dried over MgSO₄. Evaporation of the solvent in vacuo afforded the title compound as an amber oil (114.2 g, 75%).

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(b) 4-(4-Fluorophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)-1H-imidazole - To a solution of 1-(t-butylidimethylsilyloxy)-2-(4-fluorophenyl)-1-(4-pyridyl)ethanone (6.3 g, 18.3 mmol) in glacial acetic acid (125 mL) was added anhydrous copper (II) acetate (6.6 g, 36.5 mmol), ammonium acetate (14 g, 183 mmol) and 4-(methylthio)benzaldehyde (3.5 g, 22.9 mmol) and the mixture was heated at reflux. After 1 hr, the reaction was cooled then poured into a mixture of conc. NH₄OH (175 mL), ice (100 mL) and EtOAc (100 mL). The resulting mixture was stirred for 15 min, then the layers were separated. The aqueous layer was extracted with EtOAc (2 x 50 mL). The combined EtOAc extracts were washed with saturated NaCl and dried over MgSO₄. Evaporation of solvent in vacuo gave an oil which was taken up in acetone. 3 N HCl was added dropwise to adjust the pH to 2-3, and the resulting solid was filtered. The title compound [also prepared in Ex. 17 as the free base] was isolated as the yellow hydrochloride salt (3.7 g, 51%).

Example 80 - 2-[4-{(N-Benzyl-N-methyl) aminomethyl}phenyl]4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole

(a) 4-[(N-benzyl-N-methyl)aminomethyl]benzaldehyde diethylacetal - To 62.4 g (0.30 mol) of terephthalaldehyde monodiethyl acetal was added 32.1 g (0.30 mol) of benzyl amine and 500 mL toluene. The resulting solution was heated at reflux using a Dean-Stark trap. After 1 hour the solution was cooled and concentrated to give a light yellow oil (89.1 g). The oil was dissolved in 900 mL of EtOAc and 2.0 g of 5% palladium on charcoal was added. The mixture was hydrogenated on a Parr hydrogenation apparatus under 37 psi hydrogen pressure. The mixture was shaked for 1 hour at rt. The bottle was vented and 34.4 mL (0.42 mol) of 37.5% formaldehyde solution (aqueous) was added. The bottle was repressurized with 33 psi hydrogen and the mixture was shaken for 17 hours at rt. The bottle was vented and the reaction mixture was filtered and the filtrate concentrated to a nearly colorless oil (93.9 g). Vacuum distillation gave 71.4 g (76%) of 4-(N-methyl-N-benzyl)aminomethylbenzaldehyde diethylacetal: bp (30 torr) 212-234 °C.

(b) 2-[4-{(N-Benzyl-N-methyl) aminomethyl}phenyl]4-(4-fluorophenyl)-5-(4-pyridyl)-1H-imidazole - The title compound [also prepared in Ex. 36] was prepared as described in Example 13, except using 4-[(N-benzyl-N-methyl)aminomethyl]benzaldehyde diethylacetal.

**BIOLOGICAL EXAMPLES**

The cytokine-inhibiting effects of compounds of the present invention were determined by the following *in vitro* assays:
1. **IL-1** - Human peripheral blood monocytes were isolated and purified from either fresh blood preparations from volunteer donors, or from blood bank buffy coats, according to the procedure of Colotta *et al.*, *J Immunol.*, 122, 936 (1984). These monocytes (1x10^6) were plated in 24-well plates at a concentration of 1-2 million/ml per well. The cells were allowed to adhere for 2 hours, after which time non-adherent cells were removed by gentle washing. Test compounds were then added to the cells for 1h before the addition of lipopolysaccharide (50 ng/ml), and the cultures were incubated at 37°C for an additional 24h. At the end of this period, culture supernatants were removed and clarified of cells and all debris. Culture supernatants were then immediately assayed for IL-1 biological activity, either by the method of Simon *et al.*, *J. Immunol.* Methods, 84, 85, (1985) (based on ability of IL-1 to stimulate a Interleukin 2 producing cell line (EL-4) to secrete IL-2, in concert with A23187 ionophore) or the method of Lee *et al.*, *J. Immunotheraphy*, 6 (1), 1-12 (1990) (ELISA assay). Compounds of formula (I) were shown to be inhibitors of *in vitro* IL-1 produced by human monocytes.

2. **TNF** - Human peripheral blood monocytes were isolated and purified from either blood bank buffy coats or plateletpheresis residues, according to the procedure of Colotta, R. *et al.*, *J Immunol.*, 122(2), 936 (1984). The monocytes were plated at a density of 1x10^6 cells/ml medium/well in 24-well multi-dishes. The cells were allowed to adhere for 1 hour after which time the supernatant was aspirated and fresh medium (1ml, RPMI-1640, Whitaker Biomedical Products, Whitaker, CA) containing 1% fetal calf serum plus penicillin and streptomycin (10 units/ml) added. The cells were incubated for 45 minutes in the presence or absence of a test compound at 1nM-10μM dose ranges (compounds were solubilized in dimethyl sulfoxide/ethanol, such that the final solvent concentration in the culture medium was 0.5% dimethyl sulfoxide/0.5% ethanol). Bacterial lipopolysaccharide (*E. coli* 055:B5 [LPS] from Sigma Chemicals Co.) was then added (100 ng/ml in 10 ml phosphate buffered saline) and cultures incubated for 16-18 hours at 37°C in a 5% CO₂ incubator. At the end of the incubation period, culture supernatants were removed from the cells, centrifuged at 3000 rpm to remove cell debris. The supernatant was then assayed for TNF activity using either a radio-immuno or an ELISA assay, as described in WO 92/10190 and by Becker *et al.*, *J Immunol.*, 1991, 147, 4307. Compounds of formula (I) were shown to be inhibitors of *in vitro* TNF production.

IL-1 and TNF inhibitory activity does not seem to correlate with the property of the compounds of Formula (I) in mediating arachidonic acid
metabolism inhibition, further the ability to inhibit production of prostaglandin and/or leukotriene synthesis, by nonsteroidal anti-inflammatory drugs with potent cyclooxygenase and/or lipoxygenase inhibitory activity does not mean that the compound will necessarily also inhibit TNF or IL-1 production, at non-toxic doses.

3. **IL-8** - Primary human umbilical cord endothelial cells (HUVEC) (Cell Systems, Kirland, Wa) were maintained in culture medium supplemented with 15% fetal bovine serum and 1% CS-HBGF consisting of aFGF and heparin. The cells were then diluted 20-fold before being plated (250μl) into gelatin coated 96-well plates. Prior to use, culture medium was replaced with fresh medium (200μl). Buffer or test compound (25μl, at concentrations between 1 and 10μM) was then added to each well in quadruplicate wells and the plates incubated for 6h in a humidified incubator at 37°C in an atmosphere of 5% CO₂. At the end of the incubation period, supernatant was removed and assayed for IL-8 concentration using an IL-8 ELISA kit obtained from R&D Systems (Minneapolis, MN). All data were presented as mean value (ng/ml) of multiple samples based on the standard curve. IC₅₀'s where appropriate were generated by non-linear regression analysis. The compounds of formula (I), examples 5, 8b and 9, demonstrated a dose dependent reduction in the production of IL-8 (a 50-65% inhibition of IL-8).
Claims

1. A compound of formula (I):

   R1   R2
   ||
   N   > N
   R3

   (I)

   wherein
   
   R1 is 4-pyridyl, pyrimidinyl, quinolyl, isoquinoliny1, 1-imidazolyl or 1-
   benzimidazolyl which is unsubstituted or substituted with one or
   two substituents each of which is independently selected from C1-4
   alkyl, halo, C1-4 alkoxy, C1-4 alkylthio, NH2, mono- or di-C1-6-
   alkylamino or N-heterocyclyl ring which ring has from 5 to 7
   members and optionally contains an additional heteroatom
   selected from oxygen, sulfur or NR22;
   
   R2 is R8 or -OR12;
   R3 is -XaP(Z)(XbR13)2 or an optionally substituted aryl or heteroaryl
   group Q;
   
   Xa is -NR8-, -O-, -S- or a C1-10 alkylene chain optionally substituted by
   C1-4 alkyl and optionally interrupted by -NR8-, -O- or -S-;
   
   Xb is -(CR10R20)n, -NR8-, -O- or -S-;
   
   Z is oxygen or sulfur;
   
   n is 0 or an integer from 1 to 10;
   
   R4 is phenyl, naphth-1-yl or naphth-2-yl which is optionally
   substituted by one or two substituents, each of which is
   independently selected, and which, for a 4-phenyl, 4-naphth-1-yl or
   5-naphth-2-yl substituent, is halo, cyano, C(Z)NR7R17, C(Z)OR23,
   -(CR10R20)mCOR36, -SR5, -SOR5, -OR36, halo-substituted-C1-4 alkyl,
   C1-4 alkyl, -ZC(Z)R36, -NR10C(Z)R23, or -(CR10R20)mNR10R20 and
   which, for other positions of substitution, is halo, cyano,
   C(Z)NR16R26, C(Z)OR8, -(CR10R20)mCOR8, S(O)mR8, OR8, halo-
   substituted-C1-4 alkyl, -C1-4 alkyl, -NR10C(Z)R8, -NR10S(O)mR11,
   -ZC(Z)R8 or -(CR10R20)mNR16R26;
   
   m is 0, 1 or 2;
   
   R5 is hydrogen, C1-4 alkyl, C2-4 alkenyl, C2-4 alkynyl or NR7R17,
   excluding -SR5 being -SNR7R17 and -SOR5 being -SOH or ;
   
   R6 is C1-4 alkyl, halo-substituted-C1-4 alkyl, C2-4 alkenyl, C2-4 alkynyl
   or C3-5 cycloalkyl;
R₇ and R₁₇ is each independently selected from hydrogen or C₁-₄ alkyl or R₇ and R₁₇ together with the nitrogen to which they are attached form a heterocyclic ring of 5 to 7 members which ring optionally contains an additional heteroatom selected from oxygen, sulfur or NR₂₂;

R₅ is hydrogen, heterocycl, heterocyclalkyl or R₁₁;
R₁₀ and R₂₀ is each independently selected from hydrogen or C₁-₄ alkyl;
R₁₁ is C₁-₁₀ alkyl, halo-substituted C₁-₁₀ alkyl, C₂-₁₀ alkenyl, C₂-₁₀ alkynyl, C₃-₇ cycloalkyl, C₅-₇ cycloalkenyl, aryl, arylalkyl, heteroaryl or heteroarylalkyl;
R₁₂ is hydrogen, -C(Z)R₁₃ or optionally substituted C₁-₄ alkyl, aryl or aryl-C₁-₄ alkyl;
R₁₃ is hydrogen, C₁-₁₀ alkyl, cycloalkyl, heterocycl, aryl, arylalkyl, heteroaryl or heteroarylalkyl;
R₁₆ and R₂₆ is each independently selected from hydrogen or optionally substituted C₁-₄ alkyl, aryl or aryl-C₁-₄ alkyl, or together with the nitrogen which they are attached form a heterocyclic ring of 5 to 7 members which ring optionally contains an additional heteroatom selected from oxygen, sulfur or NR₁₂;
R₂₂ is R₁₀ or C(Z)-C₁-₄ alkyl;
R₂₃ is C₁-₄ alkyl, halo substituted C₁-₄ alkyl, or C₃-₅ cycloalkyl;
R₃₆ is hydrogen or R₂₃;
or a pharmaceutically acceptable salt thereof;
and excluding 2-(4-chlorophenyl)-4-(4-methoxyphenyl)-5-(4-pyridyl)imidazole.

2. A compound as claimed in claim 1 wherein R₁ is optionally substituted 4-pyridyl, 4-pyrimidinyl, 4-quinolyl, 6-isoquinolinyln, 1-imidazolyl or 1-benzimidazolyl.

3. A compound as claimed in claim 1 or 2 wherein R₂ is hydrogen or C₁-₁₀ alkyl.

4. A compound as claimed in any one of claims 1 to 3 wherein, in R₃, the group Q comprises an optionally substituted phenyl, pyrrolyl, pyridyl or pyrimidyl moiety.
5. A compound as claimed in any one of claims 1 to 4 wherein Q is substituted by up to three substituents Y1 each of which is independently selected from C1-5 alkyl, halo-substituted C1-5 alkyl, halogen, -Xp-P(Z)-(XqR13)2 or -(CR10R20)nY2 wherein Y2 is -OR8, -NO2, -S(O)mR11, -SR8, -S(O)mOR8, -S(O)mNR8R9, -NR8R9, -O(CR10R20)nNR8R9, -C(O)R8, -CO2R8, -CO2(CR10R20)nCONR8R9, -ZC(O)R8, -CN, -C(Z)NR8R9, -NR10C(Z)R8, -C(Z)NR8OR9, -NR10C(Z)NR8R9, -NR10S(O)mR11, -N(O(R21)C(Z)NR8R9, -N(O(R21)C(Z)R8, -C(=NOR21)R8, -NR10C(=NR15)SR11, -NR10C(=CR14R24)SR11, -NR10C(=CR14R24)NR8R9, -NR10C(O)(C)(O)NR8R9, -NR10C(O)(C)O)OR10, -C(=NR13)NR8R9, -C(=NOR13)NR8R9, -C(=NR13)ZR11, -OC(Z)NR8R9, -NR10S(O)mCF3, -NR10C(Z)OR10, 5-(R18)-1,2,4-oxadiazol-3-yl or 4-(R12)-5-(R18)R19)-4,5-dihydro-1,2,4-oxadiazol-3-yl; m is 1 or 2; n is 1 to 10; R9 is hydrogen, C1-10 alkyl, C2-10 alkenyl, C2-10 alkynyl, C3-7 cycloalkyl, C5-7 cycloalkenyl, aryl, arylalkyl, heteroaryl or heteroaryalkyl or R8 and R9 may together with the nitrogen to which they are attached form a heterocyclic ring of 5 to 7 members which ring optionally contains an additional heteroatom selected from oxygen, sulfur or nitrogen, selected from halogen, C1-4 alkyl, C3-7 cycloalkyl or aryly; R15 is hydrogen, cyano, C1-4 alkyl, C3-7 cycloalkyl or aryl; R18 and R19 is each independently selected from halogen, C1-4 alkyl, substituted alkyl, optionally substituted aryl, optionally substituted aryalkyl or together denote oxygen or sulfur; and R21 is hydrogen, a pharmaceutically acceptable cation, alkyl, cycloalkyl, aryl, arylC1-4alkyl, heteroaryl, heteroaryllC1-4alkyl, heterocyclyl, aryl, C1-10alkoyl.

6. A compound as claimed in claim 5 wherein the substituent Y1 is selected from halogen, C1-5 alkyl and -(CR10R20)nY2 wherein Y2 is -OR8, -NO2, -S(O)mR11, -SR8, -S(O)mNR8R9, -NR8R9, -O(CR10R20)nNR8R9, -C(O)R8, -CO2R8, -CO2(CR10R20)nCONR8R9, -CN, -C(Z)NR8R9, -NR10S(O)mR11, -NR10C(Z)R8, -NR10C(Z)NR8R9, -N(O(R21)C(Z)NR8R9, -C(=NOR21)R8, -NR10C(=NR15)SR11, -NR10C(=CR14R24)SR11, -NR10C(=CR14R24)NR8R9, -NR10C(O)(C)(O)NR8R9, -NR10C(O)(C)O)OR10, -C(=NR13)NR8R9, -C(=NOR13)NR8R9, -C(=NR13)ZR11, -OC(Z)NR8R9, -NR10S(O)mCF3, -NR10C(Z)OR10, 5-(R18)-1,2,4-oxadiazol-3-yl or 4-(R12)-5-(R18)R19)-4,5-dihydro-1,2,4-oxadiazol-3-yl.

7. A compound as claimed in claim 6 wherein the group Q has one substituent Y1 which is selected from -(CR10R20)nY2 wherein: n is 0,
1, 2 or 3, and Y₂ is -OR₈, -NO₂, -S(O)ₓR₁₁, -SR₈, -S(O)ₓNR₉R₉,
-NR₉R₉, -O(CR₁₀R₂₀)ₓNR₉R₉, -C(O)R₈, -CO₂R₈,
-CO₂(CR₁₀R₂₀)ₓCONR₉R₉, -CN, -C(Z)NR₉R₉, -NR₁₀S(O)ₓR₁₁,
-NR₁₀C(Z)R₉, -C(Z)NR₉OR₉, -NR₁₀C(Z)NR₉R₉, -N(O)R₂₁C(Z)NR₉R₉,
-C(=NOR₁₃)NR₉R₉, -NR₁₀C(=NR₁₅)NR₉R₉, 5-(R₁₈)-1,2,4-oxadiazole-3-yl
and 4-(R₁₂)-5-(R₁₈R₁₉)-4,5-dihydro-1,2,4-oxadiazol-3-yl.

8. A compound as claimed in claim 7 wherein, in Y₁, n is 0 or 1 and
Y₂ is -OH, -S(O)ₓR₁₁, -SR₈, -NR₉R₉, -CO₂R₈, -S(O)ₓNR₉R₉,
-NR₁₀S(O)ₓR₁₁, 5-(R₁₈)-1,2,4-oxadiazole-3-yl or 4-(R₁₂)-5-(R₁₈R₁₉)-4,5-
dihydro-1,2,4-oxadiazol-3-yl.

9. A compound as claimed in any one of claims 1 to 8 wherein, in R₄,
a 4-phenyl, 4-naphth-1-yl or 5-naphth-2-yl substituent is halogen,
-SR₅, -SOR₅, -OR₃₆, -(CR₁₀R₂₀)ₓNR₁₀R₂₆, or -NR₁₀C(Z)R₈ and a
substituent for another position of substitution is selected from
halogen, -S(O)ₓR₈, -OR₈, -(CR₁₀R₂₀)ₓNR₁₀R₂₆, -NR₁₀C(Z)R₈ or
-NR₁₀S(O)ₓR₁₁.

10. A compound as claimed in any one of claims 1 to 9 wherein R₁ is
4-pyridyl, 2-alkyl-4-pyridyl or 4-quinolyl; R₂ is hydrogen or methyl; R₃
is phenyl or phenyl substituted, preferably at the 4-position, with a
substituent selected from -(CR₁₀R₂₀)ₓY₂ wherein Y₂ is wherein n is 0,
1 2 or 3 and Y₂ is -OR₈, -NO₂, -S(O)ₓR₁₁, -SR₈, -S(O)ₓNR₉R₉,
-NR₉R₉, -O(CR₁₀R₂₀)ₓNR₉R₉, -C(O)R₈, -CO₂R₈,
-CO₂(CR₁₀R₂₀)ₓCONR₉R₉, -CN, -C(Z)NR₉R₉, -C(Z)NR₉OR₉,
-NR₁₀S(O)ₓR₁₁, -NR₁₀C(Z)R₈, -NR₁₀C(Z)NR₉R₉, -C(=NOR₁₃)NR₉R₉,
-NR₁₀C(=CR₁₄R₂₄)NR₉R₉, 5-(R₁₈)-1,2,4-oxadiazole-3-yl, 4-(R₁₂)-5-
(R₁₈R₁₉)-4,5-dihydro-1,2,4-oxadiazol-3-yl or a 3,5-dimethyl- or 3,5-
dibromo-4-hydroxy grouping; and R₄ is phenyl or phenyl substituted
at the 4-position with fluoro and/or substituted at the 3-position with
fluoro, chloro, C₁₋₄ alkoxy, methane-sulfonamido or acetamido.

11. A compound of formula (I) as defined in claim 1 selected from:
4-(4-Fluorophenyl)-2-(4-methylthiophenyl)-5-(4-pyridyl)imidazole;
4-(4-Fluorophenyl)-2-(4-ethylthiophenyl)-5-(4-pyridyl)imidazole;
4-(4-Fluorophenyl)-2-(4-methylsulfonylethylphenyl)-5-(4-pyridyl)imidazole;
4-(4-Fluorophenyl)-2-(4-methylnaphthylphenyl)-5-(4-pyridyl)imidazole;
4-(4-Fluorophenyl)-2-(4-ethylsulfanylphenyl)-5-(4-pyridyl)imidazole;  
4-(3-Chlorophenyl)-2-(4-methylsulfanylphenyl)-5-(4-pyridyl)imidazole;  
2-[4-(N-Methyl-N-benzyl)aminomethylphenyl]-4-(4-fluorophenyl)-5-(4-pyridyl)imidazole;  
4-(4-Fluorophenyl)-5-[4-(2-methylpyridyl)]-2-(4-methylthiophenyl)imidazole;  
4-(4-Fluorophenyl)-5-[4-(2-methylpyridyl)]-2-(4-methylsulfanylphenyl)imidazole;  
4-(4-Fluorophenyl)-2-(4-methylsulfanylphenyl)-5-(4-quinolyl)imidazole;  
2-[4-(N-Morpholino)methylphenyl]-4-(4-fluorophenyl)-5-(4-pyridyl)imidazole; or  
a pharmaceutically acceptable salt thereof.

12. A process for preparing a compound of formula (I) as defined in  
any one of claims 1 to 11 which process comprises:  
(i) condensing an α-diketone of formula (II):  
\[ R_1COCOR_4 \]  
(II)  
wherein \( R_1 \) and \( R_4 \) are as hereinbefore defined, or an equivalent  
thereof,  
with an aldehyde of the formula (III):  
\[ R_3CHO \]  
(III)  
wherein \( R_3 \) is as hereinbefore defined, or an equivalent thereof, and, if  
necessary, with ammonia or a source thereof, under imidazole-ring  
forming conditions;  
(ii) treating an α-hydroxyketone compound of formula (IIA):  
\[ R'CHOHOCOR'' \]  
(IIA)  
wherein one of \( R' \) and \( R'' \) is \( R_1 \) and the other is \( R_4 \), a suitably protected  
derivative thereof or the α-hydroxy-oxime or α-halo ketone derivative  
derivative thereof, with an oxidising agent capable of converting said compound  
into the corresponding α-diketone, in the presence of an aldehyde of  
formula (III) or an equivalent thereof, and a source of ammonia;  
(iii) treating an amidine of formula (IV):  
\[ R_3C(=NH)NHR_2 \]  
(IV)  
wherein \( R_2 \) and \( R_3 \) are as hereinbefore defined, or a salt thereof, with  
a reactive ester of an α-hydroxyketone of formula (IIA) or the  
corresponding α-haloketone, in an inert solvent, at a moderately  
elevated temperature and, if necessary, in the presence of a suitable  
condensation agent;  
(iv) treating an iminoether of formula (V):  
-64-
R₃C=NOR \hspace{1cm} (V)

wherein R₃ is as hereinbefore defined and R is C₁-₁₀ alkyl, aryl or aryl C₁-₄ alkyl, with an α-aminoketone of the formula (VI):

R'CHNH₂COR'' \hspace{1cm} (VI)

wherein one of R' and R'' is R₁ and the other is R₄ in a suitable solvent;

(v) treating the anion of an amide of formula (VII):

R₁CH₂NR₂COR₃ \hspace{1cm} (VII)

wherein R₁ and R₃ are as hereinbefore defined and R₂ is as hereinbefore defined other than hydrogen, with:

(a) a nitrile of the formula (VIII):

R₄CN \hspace{1cm} (VIII)

wherein R₄ is as hereinbefore defined, or

(b) an excess of an acyl halide, of the formula (IX):

R₄COHal \hspace{1cm} (IX)

wherein R₄ is as hereinbefore defined and Hal is halogen, or a corresponding anhydride, to give a bis-acylated intermediate which is then treated with a source of ammonia;

(vi) treating a compound of formula (X):

R'COCHR"X₇COR₃ \hspace{1cm} (X)

wherein R', R" and R₄ are as hereinbefore defined and X is O or NH, with a source of ammonia, or cyclising the corresponding Schiff's base, formed by treating the compound of formula (X) with an amine R₂NH₂;

(vii) coupling a suitable derivative of a compound of formula (XI):

```
  T₁ \   \ T₂
  \   /   \\
  N   T₃
  \  /    \\
 T₄
```

(XI)

wherein: T₂ is a nitrogen protecting group or R₂, other than hydrogen; and T₁ is hydrogen, T₃ is Q and T₄ is R₄; T₁ is R₁, T₃ is hydrogen and T₄ is R₄; or T₁ is R₁, T₃ is Q and T₄ is hydrogen, in which R₁, R₂, R₃ , R₄ and Q are as hereinbefore defined; with: (i) when T₁ is hydrogen, a suitable derivative of the heteroaryl ring R₁H, under ring coupling conditions, to effect coupling of the heteroaryl ring R₁ to the imidazole nucleus at position 5; (ii) when T₃ is hydrogen, a suitable derivative of the aryl or heteroaryl ring QH,
under ring coupling conditions, to effect coupling of the ring Q to the
imidazole nucleus at position 2; or (iii) when T₄ is hydrogen, a
suitable derivative of the aryl ring R₄H, under ring coupling
conditions, to effect coupling of the aryl ring R₄ to the imidazole
nucleus at position 4;
(viii) treating a compound of formula (XI), wherein T₁ is
hydrogen, with an N-acyl heteroaryl salt, to give an intermediate in
which the heteroaryl ring is attached to the imidazole nucleus and is
present as a 1,4-dihydro derivative thereof, which intermediate is then
subjected to oxidative-deacylation conditions; and
thereafter and if necessary carrying out all or any of the
additional steps of removing a protecting group, transforming an
initially obtained compound of formula (I) into a further compound of
formula (I) or forming a pharmaceutically acceptable salt.

13. A compound of formula (I), as defined in any one of claims 1 to 11,
or a pharmaceutically acceptable salt thereof, for use in therapy.

14. The use of a compound of formula (I), as defined in any one of
claims 1 to 11, or a pharmaceutically acceptable salt thereof, in the
manufacture of a medicament for treating a cytokine-mediated
disease state.

15. A pharmaceutical composition comprising an effective, non-toxic
amount of a compound of formula (I), as defined in any one of claims
1 to 11, or a pharmaceutically acceptable salt thereof, and a
pharmaceutically acceptable carrier or diluent.
**INTERNATIONAL SEARCH REPORT**

**I. CLASSIFICATION OF SUBJECT MATTER**

According to International Patent Classification (IPC) or to both National Classification and IPC

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**II. FIELDS SEARCHED**

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**III. DOCUMENTS CONSIDERED TO BE RELEVANT**

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<td>DE, A, 2 221 546 (CIBA-GEIGY A.G.) 16 November 1972 see claims 1,31-33, 36, 38, 43, 49, 51 see claims 52, 61-64 &amp; US, A, 3 929 807 cited in the application</td>
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**IV. CERTIFICATION**

Date of the Actual Completion of the International Search: 14 JUNE 1993

Date of Mailing of this International Search Report: 21. 06. 93

International Searching Authority: EUROPEAN PATENT OFFICE

Signature of Authorized Officer: HENRY J.C.
### ANNEX TO THE INTERNATIONAL SEARCH REPORT

**ON INTERNATIONAL PATENT APPLICATION NO.**

**US 9300674**

**SA 70064**

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 14/06/93. The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

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