MONITOR was molecules

cholesterol and inhibitors of this enzyme therefore have potenial as antihypercholesteremic agents. Turabi, N. and coworkers [*Bioorg. Med. Chem.* (1995) 3, 1479–1484] report the synthesis and biological evaluation of a series of oxime ether analogs of pravastatin, a known potent antihypercholesteremic agent. Compound **6** was found to be several times more potent than pravastatin.

α₂-Adrenoceptor antagonists

Presynaptic α_2 -adrenoceptors modulate the release of noradrenaline from nerve endings through a local feedback mechanism. α₂-Adrenoceptor antagonists should therefore increase synaptic concentrations of noradrenaline and the resultant postsynaptic stimulation of α- and β-adrenoceptors. Such agents may have clinical utility in the treatment of diabetes, depression and male sexual dysfunction. Kitchin, J. and coworkers [Bioorg. Med. Chem. (1995) 3, 1595-1603] report the synthesis and evaluation of a series of tetrahydrobenzodioxinopyrroles, such as 7, as potent, selective α_2 -adrenoceptor antagonists.

In the same journal, Beeley, L.J. and coworkers [*Bioorg. Med. Chem.* (1995) 3, 1693-1698] describe the chiral synthesis of R-(–)-2,3-dihydroisoindolylmethylimidazole (BRL 48962, **8**), a potent, selective α_{2A} -adrenoceptor antagonist, and the characterization of

this compound using cloned human α -adrenoceptors.

These studies demonstrated that the compound was 30 times more selective for the α_{2A} -adrenoceptor subtype.

Thromboxanemodulating agents

The use of thromboxane A2 (TxA2) synthase inhibitors to prevent the vasoconstrictive and platelet aggregatory actions of TxA, have been of limited success because the biosynthetic precursor, prostaglandin H2 (PGH2), is also a potent TxA, receptor agonist. TxA, receptor antagonists offer an alternative because they block the action of both TxA, and PGH2. However, such agents do not have the potential advantage of the TxA, synthase inhibitors, in that the accumulated PGH, may be utilized by PGI, synthase to produce the vasodilator and anti-aggregatory PGI2. Dickenson, R.P., Dack, K.N. and Steele, J. [Bioorg. Med. Chem. Lett. (1995) 5, 3017-3022] have therefore designed a series of dual thromboxane synthase inhibitor/thromboxane receptor antagonists by combining the key structural features for each activity in a single molecule.

This approach has yielded a number

of indole-5-propanoic acid derivatives **9**, which are potent dual agents *in vitro*.

15-Lipoxygenase inhibitor

A mass screening of the Parke-Davis compound portfolio has identified **10** as a potent 15-lipoxygenase inhibitor [Tait, B.D. *et al. Bioorg. Med. Chem. Lett.* (1996) 6, 93–96].

This compound will facilitate an investigation into the role of 15-lipoxygenase in atherosclerosis.

HIV-1 inhibitor

Navé J.F. and coworkers [Bioorg. Med. Chem. Lett. (1996) 6, 179–184] report the synthesis, phosphorylation by guanylate kinase, anti-HIV-1 and anti-herpes virus activity of two acyclic dienyl phosphonate derivatives of guanine.

Compound 11 was found to be phosphorylated by guanylate kinase and to be a significant inhibitor of HIV-1 replication. This work illustrates the potential for utilizing the dienyl phosphonate as a substitute for the phosphate group employed in other acyclonucleotide analogues.

HIV-1 integrase inhibitor

Recent attention has been directed toward HIV-1 integrase as an alternative therapeutic target for anti-HIV therapy. This protein mediates the integration of the viral DNA into the host genome and is therefore essential for survival of the virus. Eich, E. and coworkers [*J. Med. Chem.* (1996) 39, 86–95] describe the selection of the lignanolide (–)-arcrigenin **12** as a lead structure for the development of inhibitors of HIV-1 integrase.

Structure-activity studies utilizing natural, semisynthetic and synthetic lignands demonstrated that the biological activity is dependent on (i) the presence of the lactone moiety and the number and (ii) the arrangement of the phenolic hydroxyl groups on the molecule.