Aziridination of cyclic dienes with enantiopure 3-acetoxyaminoquinazolin-4(3H)-ones

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Aziridination of cyclopentadiene and cyclohepta-1,3-diene with (S)-3-acetoxyamino-2-(3-hydroxy-2,2-dimethylpropyl)quinazolin-4(3H)-one 6 (Q¹NHOAc) in the presence of titanium(IV) tert-butoxide in dichloromethane takes place highly diastereoselectively: X-ray structure determinations show that the preferred sense of diastereoselectivity in both cases is the same as that previously found for aziridination of butadiene with 6. Aziridination of cyclohexa-1.3-diene with 6 was less diastereoselective in dichloromethane solution but highly diastereoselective in acetonitrile: in this solvent two diastereoisomeric cis-4-(Q¹-amino)cyclohexen-3-ols 27 and 28 were also obtained as by-products. The same two amino alcohols were obtained by ring-opening of the aziridine with acid and were each converted into Q¹-free oxazolidinones having optical rotations which were similar in magnitude but opposite in sign.

Enantiopure cyclic vinylaziridines 1 are potentially useful relay compounds for the synthesis of a range of 1,2,3,4-tetrasubstituted cyclic amine derivatives of defined configuration and (n + 1)-azabicyclo[n.1.0] derivatives 2 by vinylaziridine pyrroline rearrangement. Because of the dearth of methods for stereoselective aziridination,^{2,3} the most direct route to these compounds 1 from the corresponding dienes (Scheme 1) has not

(CH₂)_n (enantiopure aziridinating agent)
$$\mathbf{1}$$
 (enantiopure) $\mathbf{2}$

Scheme 1

been explored. Consequently, presently available methods for synthesis of enantiopure 1 start with enantiopure dienes (substrate-controlled diastereoselectivity) and are therefore less general.4

3-Acetoxyaminoquinazolinones 3 (QNHOAc) convert alkenes into aziridines in a reaction which resembles the conversion of alkenes into epoxides by peroxyacetic acid 4 (Scheme 2).2 In both cases, 3-membered ring formation takes

$$R^{(*)}$$
 NHOAc 4 $X = NQ$ aziridination $X = O$ epoxidation

Scheme 2

place stereospecifically with retention of alkene configuration in the product. However, one advantage in the use of QNHOAc 3 is that high or complete diastereoselectivity is possible using prochiral alkenes when the R group is chiral (R*) (reagentcontrolled diastereoselectivity).

Thus Q¹NHOAc 6, prepared in situ by N-acetoxylation of the (S)-tert-leucine-derived 5 using lead tetraacetate (LTA), aziridinates e.g. butadiene or styrene highly diastereoselectively in the presence of titanium(IV) tert-butoxide (TTB) via a transition state (TS#) geometry believed to resemble that in 7 (Scheme 3).5

endo-Overlap of the conjugated double bond in butadiene/ styrene with Q¹ leads to cis-1,2-disubstituted aziridines 8a or 9a as the kinetically-formed products: these then spontaneously N-invert to the more stable trans-1,2-disubstituted **8b** or **9b**.

The value of these aziridines lies in their potential for conversion to Q¹-free chirons by aziridine ring-opening and Q¹-N bond cleavage. Thus the styrene-derived aziridine 9b has been converted into either of the Boc-protected 1,2-diamines 10 and enant. 10 (Scheme 4).6

The high diastereoselectivity obtained in the aziridination of butadiene using Q¹NHOAc 6 in the presence of TTB encouraged us to examine its use in aziridination of cyclic dienes.

Results

Aziridination of cyclopentadiene

Since the *endo*-overlap in TS[#] 7 requires an s-cis conformation for the diene, the enforced s-cis diene conformation present in cyclic dienes should make them reactive towards aziridination with QNHOAc. Reactions of cyclopentadiene with three QNHOAc compounds were first examined. With Q2NHOAc 11, aziridine 12 (42%) was obtained as a crystalline solid: the presence of hexamethyldisilazane (HMDS) in this reaction scavenges acetic acid and raises the yield. Similarly, the use of Q³NHOAc 13 yielded the corresponding aziridine 14 (55%) (Scheme 5): the greater stability of Q³NHOAc 13 often leads to superior yields of aziridination products.8

Reaction of cyclopentadiene with Q¹NHOAc 6 in the presence of TTB and crystallisation of the crude product from diethyl ether gave aziridine 16a (25%) (Scheme 6). Although the yield of aziridine 16a is low, it is isolable without the need for chromatography because it constitutes the major part of the recovered product: the fate of the other Q¹NHOAc 6-derived product(s) is unknown. An X-ray crystal structure determination of 16a9 showed that the relative configuration was in agreement with the TS# model 15 (cf. 7 for butadiene). Unexpectedly, this crystal structure also showed that the quinazolinone and cyclopentene ring residue were cis in the aziridine: the Q-substituted nitrogen usually undergoes N-inversion

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Scheme 3 Reagents and conditions: i, LTA, CH₂Cl₂, -20 °C; ii, Ti(OBut)₄; iii, butadiene or styrene.

at temperatures $<-20\,^{\circ}\text{C}$ but in this case the barrier is raised sufficiently for the first-formed *N*-invertomer **16a** to be isolated.⁶

Scheme 6

When aziridine **16a** was dissolved in deuteriochloroform and warmed in an NMR tube for 30 min at 60 °C, signals from the exo-N-invertomer **16b** appeared in the NMR spectrum and a 1:1 equilibrium ratio of N-invertomers was established which was unchanged on further heating. The $endo \longrightarrow exo$ inversion of Q^1 is accompanied by a downfield shift for the olefinic protons (see Scheme 6).

Aziridination of cyclopentadiene with Q¹NHOAc 6 in the absence of TTB gave an unseparated mixture of products whose NMR spectrum suggested the presence of 16a and its diastereoisomer

Aziridines 12 and 14 had been previously assumed to be

present as their *exo-N*-invertomers but in the light of the unexpected stability of *endo-N*-invertomer **16a**, samples of each were heated in deuteriochloroform solution for 1 h at 60 °C: no changes in their NMR spectra were observed. A sample of aziridine **12**, moreover, remained unchanged on heating briefly to 200 °C so it can be concluded that *N*-inversion from $endo \rightarrow exo$ in these aziridines had already occurred in the aziridination. This conclusion was supported by the chemical shifts of the olefinic protons (δ 6.06 and 6.21 for **12**, δ 6.01 and 6.11 for **14**) which resemble closely those in the exo-N-invertomer **16b** (see Scheme 6).

In aziridine **16a**, therefore, for reasons as yet not clear, the barrier to *N*-inversion, is significantly higher than in **12** and **14** and *endo*- and *exo-N*-invertomers are of comparable stability.

The reaction of aziridine 12 with methylcuprate was briefly investigated: using methylmagnesium bromide in the presence of copper(1) bromide, a mixture of products (71%) (Scheme 7) was obtained which could not be separated. Treatment of the mixture with lead tetraacetate (LTA) gave two imines 18 and 19 (68%) in a 1:1 ratio whose separation was carried out by Kieselgel chromatography.

The presence of two olefinic protons having very similar multiplicity and coupling constants in imines 18 and 19 suggested that the constitution of each was the same. Their assignments as different *N*-invertomers were supported by the downfield shift of the olefinic proton adjacent to the imine in the NMR spectrum of 18 and the complementary downfield shift of the methylene protons in 19 brought about in each case by the presence of the neighbouring Q group. Support for these structure assignments to imines 18 and 19 came from their interconversion on heating at ~100 °C or even on standing at room temperature over several months.

It appears, therefore, that the mechanism of the cuprate addition is predominantly $S_{\rm N}2^\prime$ but is probably not very diastereoselective.

Aziridination of cyclohexa-1,3-diene

Reaction of cyclohexa-1,3-diene with Q²NHOAc 11 and with Q³NHOAc 13 in dichloromethane gave the corresponding aziridines 20 (43%) and 21 (59%) (Scheme 8). An X-ray crystal structure determination † of aziridine 21 (Fig. 1) confirmed the expected *exo*-orientation of the Q³ group. A sample of aziridine 20 was briefly heated to 200 °C and its NMR spectrum was found to be unchanged: the Q² group of this aziridine, therefore, can also be assumed to have an *exo*-orientation.

[†] CCDC reference number 160871. See http://www.rsc.org/suppdata/p1/b1/b102592a/ for crystallographic files in. cif or other electronic format.

Q² Q²NH
$$Q^{2}$$
 Q^{2} Q^{2}

Reagents and conditions: i, MeMgBr, CuBr; ii, LTA, -20 °C. Scheme 7

Reagents: i, cyclohexa-1,3-diene, HMDS; ii, cyclohexa-1,3-

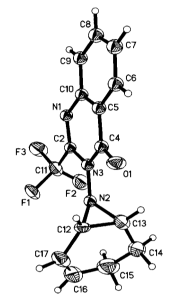


Fig. 1 Molecular structure of 21, showing the atom label scheme and 30% displacement probability ellipsoids. Hydrogen atoms are shown as spheres of arbitrary radius.

A minor product isolated from the aziridination of cyclohexa-1,3-diene with Q³NHOAc 13 was identified as the dienylamine 22 (5%).

Reaction of cyclohexa-1,3-diene with Q¹NHOAc 6 in the presence of TTB gave a mixture of aziridine 23, dienylamine 24 and tert-butoxyaminoquinazolinone 25⁵ (Scheme 9a).

Aziridine 23 was obtained as a mixture of diastereoisomers in a ratio which varied between 10:1 and 4:1 in different experiments. The major diastereoisomer was isolated as a colourless solid after crystallisation from ether-light petroleum but with considerable loss in yield. Proof of the stereostructure and the mechanism of formation of dienylamine 24 (and 22, Scheme 8) will be discussed elsewhere but in the reaction shown in Scheme 9a, dienylamine 24 was not separated from tertbutoxyaminoquinazolinone 25. Neither were the diastereoisomers of aziridine 23 separated by chromatography but their diastereoisomeric relationship was confirmed by the Swern oxidation-sodium borohydride reduction cycle in Scheme 10: a sample diastereomeric ratio (dr) 5:1 was converted to a single ketone 26 and then back to a sample of aziridine 23 of dr 3:2 showing that the two are epimeric at the aziridine ring chiral centres.

When the reaction of cyclohexa-1,3-diene with Q¹NHOAc 6 was carried out in acetonitrile as solvent, aziridine 23 was obtained in high diastereopurity although in lower yield (38%) (Scheme 9b) together with dienylamine 24 (7%) and two other products. After separation using Kieselgel chromatography, these two products were identified by NMR spectroscopy as cis-(Q1)-amino alcohols 27 and 28. In the NMR spectrum of each compound there was a coupling of ~3 Hz between the protons on adjacent carbons bearing Q¹NH and OH groups suggesting that these two protons are either both equatorial or equatorial/axial. Since it is likely that the Q¹NH group is equatorial in both compounds, the OH group will be pseudoaxial and hence the two groups are cis. Confirmation of these assignments came from conversion of each of these (Q1)-amino alcohols to the corresponding Q1-free oxazolidinone enantiomers (see below).

Higher yields of (Q1)-amino alcohols 27 and 28 were obtained by reaction of aziridine 23 (dr 5:1) with toluene-psulfonic acid in aqueous acetonitrile. After chromatography on deactivated silica, 27 and 28 were isolated in 49 and 13% yields respectively (Scheme 11). An additional product isolated in this reaction was tentatively assigned the cyclic ether structure 29: this cyclic ether was not formed from a mixture of (Q¹)-amino alcohols 27 and 28 on re-submitting it to the conditions used for the reaction shown in Scheme 11.

Some of the corresponding (Q)-amino alcohol 30 was also obtained, together with aziridine 20 and quinazolin-4(3H)-one 31 when cyclohexa-1,3-diene and Q2NHOAc 11 were reacted together in acetonitrile (Scheme 9c).

Aziridination of cyclohepta-1,3-diene and cycloheptatriene

Reaction of Q2NHOAc 11 with cyclohepta-1,3-diene gave a crystalline aziridine 32 which was isolated in 36% yield without the need for chromatography. The corresponding reaction of Q¹NHOAc 6-TTB with cyclohepta-1,3-diene gave a crude product, the major part of which was aziridine 33 (cf. aziridination of cyclopentadiene) and which crystallised directly from ethyl acetate-light petroleum in 29% yield.

An X-ray structure determination ‡ of this aziridine (Fig. 2) showed that the relative configuration was the same as that obtained for aziridine **16a** but that the Q^1 group is exo.

Aziridine 34 was obtained as a crystalline solid from the reaction of Q2NHOAc 11 and cycloheptatriene after chromatography: the other isolated product was the quinazolin-4(3*H*)-one **31** (Scheme 12b).

Discussion

Since aziridines 16a and 33 derived from cyclopentadiene and cyclohepta-1,3-diene respectively both show the same relative and absolute configuration it is likely that the major diastereoisomer from aziridination of cyclohexadiene is also formed with the same sense of diastereoselectivity.

Formation of (Q1)-amino alcohols 27, 28 and 30 (Schemes 11 and 9a) by ring-opening of the corresponding aziridines 23 and 21 with retention of configuration is suggestive of the involvement of the Q group in the reaction. Previously, ring-opening

[‡] CCDC reference number 160872. See http://www.rsc.org/suppdata/ p1/b1/b102592a/ for crystallographic files in .cif or other electronic format.

Scheme 10 Reagents: i, Swern oxidation; ii, NaBH₄, EtOH.

Scheme 11

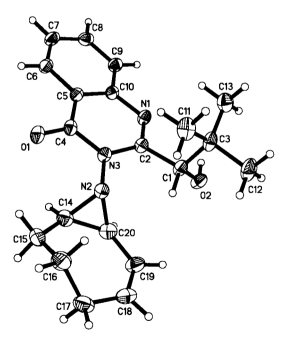


Fig. 2 Molecular structure of **33**, showing the atom label scheme and 30% displacement probability ellipsoids. Hydrogen atoms are shown as spheres of arbitrary radius.

of aziridine **8b** to the corresponding (Q¹)-amino alcohol with aqueous acetic acid was shown to involve participation of the Q¹ carbonyl oxygen.⁶

Reaction of a 4:1 mixture of aziridine diastereoisomers

$$Q^{2}NIIOAc - Q^{2}NIIOAc - Q^{2}NIIOAc - Q^{2}NIIOAc - Q^{2}NIIOAc + Q^{2}NIIOAc +$$

23 with toluene-*p*-sulfonic acid in acetonitrile containing ¹⁸O-labelled water was interrupted after 45 minutes to allow recovery of unreacted aziridine. Mass spectroscopy showed incorporation as expected of ¹⁸O label into the alcohol product but none into the recovered aziridine 23 which supports a mechanism for quinazolinone participation analogous to that proposed previously and illustrated for the reaction of the major aziridine diastereoisomer in Scheme 13.

Recovery of unlabelled aziridine 23 from this reaction serves as a control to eliminate the possibility of exchange of the (Q)C=O without its involvement in aziridine ring-opening. Much of the ¹⁸O label in the amino alcohol products 27 and 28 was lost on silicon chromatography which supports its presence in the quinazolinone carbonyl group. Scheme 13 also shows a possible competitive reaction (b in 35) of the allyl cation with the hydroxy group in the Q¹ side-chain giving cyclic ether 29.

Interestingly, the diastereoisomer ratio (dr) of the recovered aziridine in the experiment above was 10:1 and the dr of the alcohol product (27:28) was ~2:1 confirming that the minor aziridine diastereoisomer was ring-opened approximately twice as fast as the major one (and providing an expedient route for purification of the major diastereoisomer 23). In the light of these results it seems likely that the isolation of only the major aziridine diastereoisomer 23 from aziridination of cyclohexa-1,3-diene with Q¹NHOAc 6–TTB results from faster ring-opening of the minor diastereoisomer by a mechanism resembling that shown in Scheme 13 but mediated by a titanium-containing species instead of by acid.

Conversion of (Q1)-amino alcohols 27 and 28 into Q1-free chirons

One of the objectives of this work was to use the products from aziridination of dienes with enantiopure QNHOAc reagents

Scheme 13

to prepare Q-free aziridine ring-opened enantiopure products containing two or more chiral centres (chirons).

In a model reaction, Q³-amino alcohol 30 was heated in THF at reflux with sodium hydride and 1,1'-carbonyldiimidazole for 3 h. Chromatography of the crude product gave oxazolidinone 36 (53%) together with unchanged starting material 30 (35%) (Scheme 14).

30
$$\stackrel{\text{i}}{\longrightarrow}$$
 $\stackrel{\text{N}}{\longrightarrow}$ 0

36 (58%)

27 $\stackrel{\text{i}}{\longrightarrow}$ $\stackrel{\text{N}}{\longrightarrow}$ 0 $\stackrel{\text{ii}}{\longrightarrow}$ $\stackrel{\text{N}}{\longrightarrow}$ 0 $\stackrel{\text{II}}{\longrightarrow}$ 0 $\stackrel{\text{N}}{\longrightarrow}$ 0 $\stackrel{\text{II}}{\longrightarrow}$ 0 $\stackrel{\text{N}}{\longrightarrow}$ 0 $\stackrel{\text{N}}{\longrightarrow$

Scheme 14 Reagents and conditions: i, 1,1'-carbonyldiimidazole, NaH, THF, 3 h; ii, SmI₂, Bu^tOH, THF.

The similarity in rates of the corresponding reactions of (Q¹)-amino alcohols 27 and 28 with 1,1'-carbonyldiimidazole giving oxazolidinones 37 and 38 respectively supports the previously drawn conclusion that both have cis-substituted cyclohexene rings.

Previously we had shown that the presence of a carbonyl group on the exocyclic 3-aminoquinazolinone facilitated reductive cleavage of the N-N bond and allowed aluminium amalgam to be used. 10 However, reduction of oxazolidinones 37 and 38 did not take place with aluminium amalgam but was successful using samarium(II) iodide in the presence of tertbutyl alcohol.¹⁰ Q¹-free oxazolidinones 39 and enant. 39 were separated from quinazolin-4(3H)-one Q¹H 40 by chromatography and found to have optical rotations which were similar in magnitude but opposite in sign. Assignments of absolute configuration to oxazolidinones 39 and enant. 39 are based on the assumption that aziridination of cyclohexa-1,3-diene with Q¹NHOAc 6 proceeds preferentially with the same diastereosense as in aziridination of butadiene, cyclopentadiene and cyclohepta-1,3-diene.

Summary

Cyclopentadiene and cyclohepta-1,3-diene were mono-azirid-

inated highly diastereoselectively by Q1NHOAc 6 and TTB in dichloromethane and the crystalline products were isolated without the need for chromatography. Cyclohexa-1,3-diene is aziridinated highly diastereoselectively by Q1NHOAc 6 and TTB in acetonitrile and the product 23 is separated by chromatography from cis-Q1-amino alcohols 27 and 28 formed as by-products.

These enantiopure aziridines could provide a useful source of chirons since Q¹NH₂ 5, the precursor of Q¹NHOAc 6, is available from tert-leucine in 43% yield (5 steps) without the need for chromatography. Thus the two (Q1)-amino alcohols 27 and 28 have been converted into enantiomeric Q1-free oxazolidinones 39 and enant. 39.

Experimental

For details of instrumentation and other experimental details see refs. 5 and 6.

General procedure A for aziridination of dienes using Q²NHOAc 11 and Q³NHOAc 13

Dry dichloromethane (1 cm³ for each 0.1 g QNH₂) was added to a round-bottom flask suspended in a dry ice-acetone bath at -12 °C and magnetically stirred. Lead(IV) acetate (LTA; 1.05 mol equiv.) was added to the flask in one portion. When the LTA had dissolved, the temperature of the bath was lowered to -20 °C and the appropriate QNH₂ (1.0 mol equiv.) added in small portions over 10-15 min at this bath temperature, stirring throughout. The temperature of the bath was allowed to rise to −10 °C and the solution was filtered into a flask maintained at -10 °C to remove the lead diacetate produced (on a small scale, a Pasteur pipette and a cotton wool plug can be used). To this filtered solution containing the QNHOAc, the diene (1.5-3 mol equiv.) and (with Q2NH2 only) hexamethyldisilazane (3 mol equiv.) were added, the cooling bath removed and the temperature allowed to reach ambient (~15 min) stirring throughout. The reaction mixture was filtered, the filtrate was washed with saturated aqueous sodium hydrogencarbonate solution and water, dried and the solvent evaporated under reduced pressure.

General procedure B for the aziridination of dienes using Q¹NHOAc 6 in the presence of titanium(IV) tert-butoxide⁵

Procedure A was followed up to and including addition of Q¹NH₂ 5 and the cold (-10 °C) solution filtered through a cotton wool plug into a stirred solution of titanium(IV) tertbutoxide (2.1 equiv.) held at -20 °C. After stirring at this temperature for 2 min the alkene (2 equiv.) was added and the temperature of the reaction mixture allowed to rise to ambient by removal of the cooling bath. Saturated sodium hydrogencarbonate solution was added to the vigorously stirred reaction mixture and a gelatinous precipitate formed immediately. The solution was filtered through Celite and the organic layer of the filtrate was separated, washed with brine, dried, and the solvent evaporated under reduced pressure to give the crude product.

Aziridination of cyclopentadiene using Q2NHOAc 11

General aziridination procedure A was followed using 3-amino-2-isopropylquinazolin-4(3*H*)-one ¹¹ (300 mg, 1.47 mmol), LTA (687 mg, 1.55 mmol), HMDS (474 mg, 2.94 mmol) and cyclopentadiene (194 mg/0.24 cm³, 2.94 mmol) in dichloromethane (6 cm³). Crystallisation of the crude product (457 mg) from ethyl acetate-light petroleum gave aziridine 12 (159 mg, 42%) as a colourless solid, mp 126-128 °C (from ethyl acetatelight petroleum) (Found: C, 71.5; H, 6.4; N, 15.7%. C₁₆H₁₇ON₃ requires C, 71.8; H, 6.4; N, 15.7%); $v_{\text{max}}/\text{cm}^{-1}$ 1670s, 1470m and 1380s; $\delta_{\rm H}$ 1.40 (3H, d, J 6.6, CH₃CHCH₃), 1.42 (3H, d, J 6.6, CH_3CHCH_3), 2.76 (1H, dddd, J 18.9, 5.0, ~2 and ~2, CHH), 2.95 (1H, ddd, J 18.9, ~2 and ~2, CHH), 3.60 (1H, br d, J ~5 and ~5, azir. NCHCH₂), 3.67 [1H, heptet, J 6.6, CH(CH₃)₂], 3.79 (1H, br d, $J \sim 5$, azir. NCHC=C), 6.3–6.9 (1H, struct. m, CH=CH), 6.21 (1H, dddd, J 5.6, ~2, ~2, ~2, CH=CH), 7.40 [1H, ddd, J 8.2, 6.9, 1.5, H-6(Q)], 7.59–7.72 [2H, struct. m, H-7, H-8 (Q)] and 8.19 [1H, dd, J 8.2, 1.0, H-5(Q)]; $\delta_{\rm C}$ 21.6, 32.7 $(2 \times CH_3)$, 36.7 (CH₂), 49.5 [CH(CH₃)₂], 52.5, 58.1 (2 × C-N), 121.8 [CCO(Q)], 126.4, 126.5, 127.3, 128.4, 133.9, 138.5 $[4 \times CH(Q)]$ and HC=CH, 146.6 [CN=C(Q)] and 160.4, 161.8 [CN(Q), CO(Q)]; m/z(%) 267 (M⁺,10), 189 (12), 188 (43), 187 (27) and 173 (100).

Aziridination of cyclopentadiene using Q3 NHOAc 13

3-Amino-2-trifluoromethylquinazolin-4(3H)-one (Q³NH₂) was prepared by the published route⁷ except that the crude 2-trifluoromethyl-3,1-benzoxazin-4-one was treated hydrazine in ethanol directly after removal of trifluoroacetic anhydride. Following general procedure A, the foregoing Q3NH₂ (300 mg, 1.31 mmol), LTA (609 mg, 1.38 mmol) and cyclopentadiene (173 mg/0.22 cm³, 2.62 mmol) were reacted in dichloromethane (6 cm³). Crystallisation of the crude product (309 mg) from ethyl acetate-light petroleum gave aziridine 14 (212 mg, 55%) as a colourless solid, mp 125–127 °C (from ethyl acetate–light petroleum) (Found: M^+ 293.0775. $C_{14}H_{10}ON_3F_3$ requires M 293.0775); $v_{\text{max}}/\text{cm}^{-1}$ 1665s, 1470m and 1390s; δ_{H} 2.70 (1H, dddd, J 18.8, 5.0, ~2, ~2, CHH), 2.85 (1H, ddd, J 18.8, \sim 2, \sim 2, CHH), 4.04 (1H, br dd, J \sim 5, \sim 5, azir. NCHCH₂), 4.28 (1H, br d, J ~5, azir. NCHC=C), 6.01 (1H, struct. m, CH=CH), 6.11 (1H, struct. m, CH=CH), 7.59 [1H, ddd, J 8.2, ~4, ~4, H-6(Q)], 7.75–7.82 [2H, m, H-7, H-8(Q)] and 8.24 [1H, br d, J 8.2, H-5(Q)]; $\delta_{\rm C}$ 38.6 (CH₂), 49.1, 55.0 (2 × C-N), 123.3 [CCO(Q)], 127.0, 128.8, 129.0, 129.5, 134.9, 138.4 $[4 \times CH(Q)]$ and HC=CH, 144.3 [CN=C(Q)] and 160.5, 170.2 [CN(Q), CO(Q)]; m/z(%) 293 $(M^+, 7)$, 215 (100), 214 (70) and 213 (38).

Aziridination of cyclopentadiene using Q¹NHOAc 6 and TTB

General aziridination procedure **B** was followed in this reaction using Q¹NH₂ **5** (200 mg, 0.80 mmol), LTA (376 mg, 8.50 mmol), TTB (576 mg, 1.71 mmol) and cyclopentadiene (105 mg/131 cm³, 1.59 mmol) in dichloromethane (5 cm³). Crystallisation of the crude product from diethyl ether gave *endo-aziridine* **16a** (60 mg, 25%) as a colourless solid, mp 126–128 °C (from diethyl ether–light petroleum). [a]_D = +1.3 (c = 1.1, EtOH) (Found: C, 69.1; H, 6.8; N, 13.4%. C₁₈H₂₁O₂N₃ requires C, 69.4; H, 6.8; N, 13.4%); v_{max}/cm⁻¹ 1675s, 1610m and 1590s; δ _H 1.00 [9H, s, C(CH₃)₃], 2.43 (1H, ddd, J 20.1, ~5, ~2, CHH), 2.63 (1H, dddd, J 20.1, 5.5, ~2, ~2, CHH), 3.58 (1H, br dd, J ~5, ~5, azir. NCHCH₂), 3.85 (1H, d, J 10.6, CHOH), 4.19–4.22 (1H, m, azir. NCHC=C), 4.75 (1H, d, J 10.6, CHOH), 5.35–5.41 (1H, m, CH=CH), 5.65 (1H, dddd, J 5.7, ~2, ~2, ~2, CH=CH), 7.42 [1H, ddd, J 8.2, 6.9, 1.0, H-6(Q)], 7.62 [1H, dd, J 8.2, 1.0,

H-8(Q)], 7.70 [1H, ddd, J 8.2, 6.9, 1.0, H-7(Q)] and 8.17 [1H, dd, J 8.2, 1.0, H-5(Q)]; $\delta_{\rm C}$ 26.3 [C(CH₃)₃], 38.6 (C), 39.4 (CH₂), 50.6, 57.0 (2 × C-N), 75.3 (C-OH), 121.9 [CCO(Q)], 126.2, 126.6, 126.8, 127.2, 132.8, 134.0 [4 × CH(Q) and HC=CH], 145.1 [CN=C(Q)] and 156.9, 158.2 [CN(Q) and CO(Q)]; m/z(%) 311 (M⁺, 4) and 176 (100).

Thermal equilibration of *endo*-aziridine 16a and *exo*-aziridine 16b

endo-Aziridine **16a** (50 mg, 0.16 mmol) was heated in CDCl₃ at 60 °C for 30 min. On cooling, the NMR spectrum showed, in addition to the signals above belonging to *endo*-aziridine **16a**, those assignable to *exo*-aziridine **16b** at $\delta_{\rm H}$ 1.02 [9H, s, C(CH₃)₃], 2.74 (1H, dddd, *J* 18.8, ~5, ~2, ~2, C*H*H), 2.99 (1H, ddd, *J* 18.8, ~5, ~2, C*H*H), 3.63 (1H, br d, *J* ~6, azir. NC*H*C=C), 3.84 (1H, d, *J* 10.4, CHO*H*), 3.99 (1H, br dd, *J* ~5, ~5, azir. NC*H*CH₂), 5.01 (1H, d, *J* 10.4, C*H*OH), 6.07 (1H, m, C*H*=CH), 6.14 (1H, struct. m, CH=C*H*), 8.21 [1H, dd, *J* 8.2, 1.0, H-5(Q)]; $\delta_{\rm C}$ 25.9 [C(CH₃)₃], 32.5 (CH₂), 38.1 (C), 51.2, 56.6 (2 × C–N), 75.0 (C–OH), 121.5 [CCO(Q)], 125.8, 126.4, 127.0 [3 × CH(Q)], 129.0, 133.6, 139.2 [H*C*=*C*H and CH(Q)], 144.8 [*C*N=C(Q)] and 157.9, 159.6 [CN(Q), CO(Q)].

Reaction of aziridine 12 with cuprate

A flame dried 2-necked flask equipped with a septum cap and 3-way tap was flushed with nitrogen, degassed, flushed with argon, degassed and filled with argon. Using syringes, a suspension of copper(I) bromide–dimethyl sulfide (58 mg, 0.28 mmol) (prepared by the literature procedure 12) and dissolved in THF (1 cm³) was added followed by methylmagnesium bromide (101 mg/0.1 cm³, 0.84 mmol) dissolved in THF (1 cm³). Aziridine **16a** (75 mg, 0.28 mmol), dissolved in THF (1 cm³), was then added dropwise *via* a syringe to the solution which was stirred at ambient temperature for 1 h. After addition of ethyl acetate (5 cm³) the solution was washed with saturated aqueous sodium hydrogencarbonate (5 cm³) and the solvent separated, dried and evaporated. Column chromatography (3 : 1 light petroleum–ethyl acetate) of the crude product (80 mg) gave a mixture of (Q²)amines **17** (58 mg, 71%) as a yellow oil, R_f 0.28.

LTA oxidation of (Q2)amines 17

The above mixture of amines 17 (58 mg, 0.20 mmol) was dissolved in dichloromethane (1 cm³) at −20 °C and HMDS (72 mg, 0.45 mmol) was added. LTA (95 mg, 0.21 mmol) was added in small portions over 10 min and the solution allowed to warm to ambient temperature. Dichloromethane (10 cm³) was added and the solution washed with saturated aqueous sodium hydrogencarbonate (5 cm³), the organic layer separated, dried and evaporated to give the crude product as a yellow oil. Column chromatography (2:1 light petroleum-ethyl acetate) gave $N(Q^2)$ -imine 18 (18 mg, 33%) as a colourless oil, R_f 0.43 (Found: M^+ 281.1529. $C_{17}H_{19}ON_3$ requires M 281.1528); $v_{\rm max}/{\rm cm}^{-1}$ 1780 m, 1675s, 1620s and 1595 m; $\delta_{\rm H}$ 1.13 (3H, d, J 6.9, CHC H_3), 1.29 (3H, d, $J \sim 7$, CHC H_3), 1.30 (3H, d, $J \sim 7$, CHCH₃), 2.08 (1H, dd, J 18.6, ~2, CHH), 2.75 (1H, dd, J 18.6, ~6, CHH), 3.02 (1H, struct. m, CHMe), 3.31 [1H, heptet, J 6.9, $CH(CH_3)_2$, 6.55 (1H, dd, J 5.7, 2.2, CH=CHC=N), 7.01 (1H, br dd, J~6, ~2, N=CCH=CH), 7.43 [1H, ddd, J~8, ~6, ~3, H-6(Q)], 7.69-7.73 [2H, m, H-7 and H-8(Q)] and 8.29 [1H, dd, J 8.2, 1.0, H-5(Q)]; m/z (%) 281 (M⁺, 78), 266 (100), 187 (81) and 173 (73).

Further elution gave $N(Q^2)$ -imine **19** (19 mg, 35%) as a colourless oil, $R_{\rm f}$ 0.36 (Found: M⁺ 281.1529. C₁₇H₁₉ON₃ requires M 281.1528); $\nu_{\rm max}/{\rm cm}^{-1}$ 1780 m, 1675s, 1620s and 1595 m; $\delta_{\rm H}$ 1.24 (3H, d, J 6.9, CHC H_3), 1.28 (6H, d, J ~7, 2 × CHC H_3), 2.55 (1H, dd, J 18.1, ~2, CHH), 3.14 (1H, struct. m, CHMe), 3.23 (1H, dd, J 18.1, 6.9, CHH), 3.36 [1H, heptet, J 6.6, CH(CH₃)₂], 5.97 (1H, dd, J 5.7, 1.5, CH=CHC=N), 6.92 (1H, dd, J 5.7, 2.5, N=CCH=CH), 7.42 [1H, ddd, J 8.2, 6.9, 1.2,

A sample of imine 18 which had been set aside for 18 months was found to have partially interconverted with imine 19 (ratio ~2: 1 respectively).

Aziridination of cyclohexa-1,3-diene with Q²NHOAc 11

General aziridination procedure A was followed in this reaction using 3-amino-2-isopropylquinazolin-4(3H)-one 11 (600 mg, 2.96 mmol), LTA (1.38 g, 3.11 mmol), HMDS (1.19 g, 7.40 mmol) and cyclohexa-1,3-diene (470 mg/0.55 cm³, 5.92 mmol) in dichloromethane (12 cm³). Column chromatography of the crude product (0.68 g) using 2:1 light petroleum-ethyl acetate gave aziridine 20 (0.54 g, 60%) as a colourless solid, R_f 0.41, mp 93-95 °C (from light petroleum-ethyl acetate) (Found: MH+ 282.1606. $C_{17}H_{20}ON_3$ requires MH^+ 282.1607); v_{max}/cm^{-1} 1660s and 1570s; $\delta_{\rm H}$ 1.32 (3H, d, J 6.6, CH₃CHCH₃), 1.35 (3H, d, J 6.6, CH₃CHCH₃), 1.69 (1H, struct. m, CH₂CHH), 2.13 (1H, ddd, J~18, ~6, ~6, CHHCH₂), 2.27 (1H, struct. m, incl. J 18.0, CHHCH₂), 2.61 (1H, dddd, J~15, ~7, ~2, ~2, CH₂CHH), 2.94 (1H, dd, J7.9, 4.7, azir. NCHCH₂), 3.36 (1H, dd, J7.9, 2, azir. NCHC=C), 3.64 [1H, heptet, J 6.6, CH(CH₃)₂], 6.03 (1H, ddd, J 8.7, ~6, ~2, CH=CH), 6.23 (1H, ddd, J 8.7, ~6, ~4, CH=CH), 7.42 [1H, ddd, J 8.1, 6.3, 1.2, H-6(Q)], 7.60–7.70 [2H, m, H-7 and H-8(Q)] and 8.20 [1H, dd, J 8.1, 1.2, H-5(Q)]; $\delta_{\rm C}$ 20.4 (CH₂), 22.7 (CH₃), 23.1 (CH₂), 23.4 (CH₃), 33.3 [CH(CH₃)₂], 45.8, 50.8 (2 × C–N), 123.3, 123.6 [HC=CH, CCO(Q)], 128.3, 129.2, 135.5, 135.7 $[4 \times CH(Q)]$, 148.4 [CN=C(Q)] and 162.4, 163.7 [CN(Q), CO(Q)]; m/z(%) 282 (MH⁺, 100) and 189 (45).

Aziridination of cyclohexa-1,3-diene with Q3NHOAc 13

General aziridination procedure A was followed using 3-amino-2-trifluoromethylquinazolin-4(3H)-one (see above; 200 mg, 0.87 mmol), LTA (406 mg, 0.92 mmol) and cyclohexa-1,3-diene (140 mg/0.16 cm³, 1.75 mmol) in dichloromethane (4 cm³). Crystallisation of the crude product (246 mg) from ethyl acetate-light petroleum gave aziridine 21 (170 mg) as a white solid, mp 103–105 °C (Found: C, 58.6; H, 3.85; N, 13.7%. $C_{15}H_{13}ON_3F_3$ requires C, 58.6; H, 3.9; N, 13.7%); ν_{max}/cm^{-1} 1690s, 1610s, 1470m and 1380s; $\delta_{\rm H}$ 1.43 (1H, dddd, J 14.0, 10.0, 6.7, ~2, CH₂CHH), 1.91 (1H, ddd, J 17.5, ~7, ~7, CHHCH₂), 2.05 (1H, struct. m, incl. J 17.5, CHHCH₂), 2.23 (1H, dddd, J 14.0, 7.8, ~2, ~2, CH₂CHH), 3.59 (1H, dd, J 7.5, 4.8, azir. NCHCH₂), 3.82 (1H, d, J 7.5, azir. NCHC=C), 5.82 (1H, m, CH=CH), 5.97 (1H, ddd, J 9.5, ~5, ~3, CH=CH), 7.41 [1H, ddd, J 8.2, 4.9, 3.2, H-6(Q)], 7.60–7.69 [2H, m, H-7 and H-8(Q)] and 8.04 [1H, dd, J 8.0, 1.0, H-5(Q)]; δ_C 18.6, 21.2 (2 × CH₂), 38.5, 44.1 (2 × C–N), 122.1, 123.3 [CCO(Q), HC=CH], 127.2, 128.6, 129.3 [3 × CH(Q)], 133.1, 134.9 [CH(Q), HC=CH], 144.3 [CN=C(Q)], 160.9, 162.3 $[CN(Q), CO(Q)]-(CF_3 \text{ not visible})$; m/z(%) 308 (MH⁺, 100), 230 (24), 215 (46).

Kieselgel chromatography (2:1 light petroleum-ethyl acetate) of the residue after evaporation of the filtrate above gave more aziridine 21 (13 mg, total 68%), R_f 0.41.

A crystal suitable for X-ray crystallography was prepared by crystallisation from light petroleum.

Further elution gave the dienylamine 22 (13 mg, 5%) as a clear colourless oil, $R_{\rm f}$ 0.33 (Found: MH⁺ 308.1011. $C_{15}H_{13}$ - ON_3F_3 requires MH^+ 308.1011); v_{max}/cm^{-1} 1695m and 1610m; $\delta_{\rm H}$ 2.31 (1H, dddd, J ~18, ~8, ~4, ~2, CHH), 2.45 (1H, dddd, J 18.0, 5.0, 5.0, ~1.5, CHH), 3.93 (1H, m, incl. J ~11, CHNH), 5.43 (1H, d, J11, NH), 5.83 (1H, dd, J9.6, 5.1, CH=CH), 5.90 (1H, dddd, J 10.5, 5.9, ~4, ~2, CH=CH), 6.03 (1H, m, CH=CH), 6.13 (1H, dddd, J 9.6, 5.0, ~1, ~1, CH=CH), 7.64 [1H, ddd, J 8.0, ~4, ~4, H-6(Q)], 7.81-7.86 [2H, m, H-7 and H-8(Q)] and 8.31 [1H, dd, J 8.2, 1.0, H-5(Q)]; $\delta_{\rm C}$ (75 MHz) 28.3 (CH₂), 54.2 (CNH), 122.5 [CCO(Q)], 123.8, 124.0, 126.1, 127.3, 127.6, 129.2, 129.6, 135.5 [$4 \times CH(Q)$ and $2 \times HC = CH$], 145.3

[CN=C(Q)] and 162.4, 168.3 [CN(Q), CO(Q)]; m/z(%) 308 (MH⁺, 44), 307 (M⁺, 42), 230 (100) and 229 (50).

Aziridination of cyclohexa-1,3-diene with Q¹NHOAc 6-TTB

General aziridination procedure B was followed using Q¹NH₂ 5 (500 mg, 2.01 mmol), LTA (942 mg, 2.12 mmol), TTB (1.42 g, 4.20 mmol) and cyclohexa-1,3-diene (320 mg/0.38 cm³, 4.00 mmol) in dichloromethane (11 cm³). After work up, column chromatography (8:1 light petroleum-ethyl acetate) gave a mixture of dienylamine 24 (37 mg, 6%) and tert-butoxyaminoquinazolinone 25 (26 mg, 6%) as a colourless oil, $R_{\rm f}$ 0.17. For dienylamine **24**: $[a]_D = +12.9$ (c = 1.0, EtOH) (Found: M⁺ 325.1791. $C_{19}H_{23}O_2N_3$ requires M 325.1790); v_{max}/cm^{-1} 3500m, 1675s, 1595s and 1470s; $\delta_{\rm H}$ 0.96 [9H, s, (CH₃)₃], 2.37 (2H, m, CH₂), 3.61 (1H, d, J 10.4, CHOH), 3.96 (1H, ddd, J~12, ~6, ~6, CHNH), 5.13 (1H, d, J 10.4, CHOH), 5.37 (1H, d, J 5.9, NH), 5.58 (1H, br dd, J ~9, ~5, CH=CH), 5.92 (1H, m, CH=CH), 6.05 (1H, m, CH=CH), 6.15 (1H, m, CH=CH), 7.48 [1H, ddd, J 8.2, 6.9, 1.0, H-6(Q)], 7.69 [1H, d, J 8.2, H-8(Q)], 7.78 [1H, ddd, J 8.2, 6.9, 1.0, H-7(Q)] and 8.23 [1H, dd, J 8.2, 1.0, H-5(Q)]; $\delta_{\rm C}$ (75 MHz) 26.3 [(CH₃)₃], 28.3 (CH₂), 38.3 [C(CH₃)₃], 52.0 (C-N), 75.2 (COH), 120.3 [CCO(Q)], 122.9, 123.8, 126.3, 127.1, 127.2, 127.7, 128.2, 134.9 [2 \times HC=CH and $4 \times CH(Q)$], 146.9 [CN=C(Q)] and 160.6, 161.8 [CN(Q), CO(Q)]; m/z(%) 326 (MH⁺, 52), 248 (48), 233 (100) and 215 (36). 3-tert-Butoxyaminoquinazolinone 25 was identified by comparison of signals at δ 0.97 and 1.36 (lit. 5 0.91 and 1.27) in its ¹H NMR spectrum.

Further elution gave aziridine 23 (291 mg, 54%) as a colourless gum, R_f 0.11, which crystallised from diethyl ether-light petroleum, mp 106–107 °C. $[a]_D = +16.9$ (c = 2.5, EtOH) (Found: M^+ 325.1790. $C_{19}H_{23}O_2N_3$ requires M 325.1790); $v_{\rm max}/{\rm cm}^{-1}$ 3480m, 1660s and 1580s; $\delta_{\rm H}$ 1.02 [9H, s, C(CH₃)₃], 1.65 (1H, dddd, J 13.0, 12.5, 7.5, ~3, CHHCH₂), 2.15–2.20 (2H, m, CH₂), 2.66 (1H, dddd, J~13, ~8, ~2, ~2, CHHCH₂), 2.91 (1H, ddd, J7.7, 4.7, 1.0, azir. NCHCH₂), 3.57 (1H, br d, J7.7, azir. NCHC=C), 3.82 (1H, d, J 10.4, CHOH), 5.04 (1H, d, J 10.4, CHOH), 6.03 (1H, m, CH=CH), 6.22 (1H, m, CH=CH), 7.44 [1H, ddd, J 8.2, 6.9, 1.0, H-6(Q)], 7.64 [1H, d, J 8.0, H-8(Q)], 7.70 [1H, ddd, J 8.0, 6.9, 1.2, H-7(Q)] and 8.21 [1H, dd, J 8.2, 1.2, H-5(Q)]; $\delta_{\rm C}$ 18.4, 21.6 (2 × CH₂), 26.3 $[C(CH_3)_3]$, 38.5 $[C(CH_3)_3]$, 45.5, 48.7 (2 × C-N), 75.1 (COH), 120.7, 121.8 [CCO(Q), HC=CH], 126.3, 126.7, 127.0, 127.4, 134.1 [4 × CH(Q), HC=CH], 145.0 [CN=C(Q)] and 158.0, 159.9 [CN(Q), CO(Q)]; *m/z*(%) 325 (M⁺, 48), 268 (62), 240 (100), 231 (94) and 215 (80). The NMR spectrum of aziridine 23 before crystallisation showed the presence of another diastereoisomer with (observable signals)— $\delta_{\rm H}$ 0.99 [9H, s, C(CH₃)₃], 2.47 (1H, struct. m, CHH), 3.85 (1H, d, J 10.3, CHOH) and 5.02 (1H, d, J 10.3, CHOH). The ratio of aziridine diastereoisomers ranged from 10:1-4:1 in different experiments from comparison of signals at δ 2.66 and 2.47 in the ¹H NMR spectrum of the crude reaction product.

Swern oxidation of aziridine 23

DMSO (73 mg/0.07 cm³, 0.93 mmol) was added to dichloromethane (1 cm³) pre-cooled to -78 °C followed by dropwise addition of oxalyl chloride (60 mg, 0.46 mmol). After stirring for 10 min, aziridine 23 (dr 5:1) (100 mg, 0.31 mmol) was added to a solution of dichloromethane (1 cm³) and the mixture stirred at -78 °C for 2 h. Triethylamine (219 mg, 2.17 mmol) was added and the solution warmed to ambient temperature. After addition of saturated sodium hydrogencarbonate solution (10 cm³) and extraction with dichloromethane (15 cm³), drying of the organic layer and evaporation gave the crude product as a yellow oil. Column chromatography (3:1 light petroleum-ethyl acetate) gave ketone 26 (51 mg, 51%) as a colourless oil, $R_{\rm f}$ 0.50 (Found: MH⁺ 324.1712. $C_{19}H_{22}O_{2}N_{3}$ requires MH^+ 324.1712); δ_H 1.32 [9H, s, (CH₃)₃], 1.46–1.61

(1H), 2.00–2.08 (2H) and 2.30–2.39 (1H) (3 × m, CH_2CH_2), 3.69 (1H, ddd, J 7.8, 4.8, 1.5, azir. $CHCH_2$), 3.97 (1H, ddd, J 7.8, ~2, ~2, azir. CHC=C), 5.88 (1H, m, CH=CH), 6.06 (1H, m, CH=CH), 7.48 [1H, ddd, J 8.0, 6.6, 1.1, H-6(Q)], 7.66 [1H, dd, J 8.0, 1.1, H-8(Q)], 7.75 [1H, ddd, J 8.0, 6.6, 1.3, H-7(Q)] and 8.21 [1H, dd, J 8.0, 1.3, H-5(Q)]; δ_C 18.2 (CH_3)₃, 21.4, 27.1 (2 × CH_2), 40.4, 44.9, 45.0 [2 × CH_3), (1CC), 122.8 [HC=CH, CCO(Q)], 126.7, 127.6, 128.0, 132.9, 134.5 [4 × CH(Q), HC=CH], 146.2 [CN=C(Q)], 153.2, 160.3 [CN(Q), CO(Q)] and 204.9 (C=O); m/z(%) 324 (MH^+ , 31), 246 (22), 231 (100) and 215 (44).

Further elution gave unreacted aziridine 23 (19 mg, 19% recovered), R_t 0.23.

Sodium borohydride reduction of ketone 26

Ketone **26** (50 mg, 0.16 mmol) was stirred in ethanol (5 cm³) with sodium borohydride (3 mg, 0.05 mmol) at ambient temperature for 4 h. Addition of saturated sodium hydrogencarbonate solution (10 cm³) and extraction with ethyl acetate (10 cm³), drying and then evaporation of the organic layer gave the crude product as a colourless oil. Column chromatography (3:1 light petroleum–ethyl acetate) gave ketone **26** (14 mg, 28% recovered) as a colourless oil, $R_{\rm f}$ 0.51.

Further elution gave aziridine **23** (18 mg, 36%) as a colourless oil, $R_{\rm f}$ 0.30, in which the two aziridine diastereoisomers **23** were now present in a 3 : 2 ratio from comparison of signals at δ 3.57 and 3.85 in its ¹H NMR spectrum.

Aziridination of cyclohepta-1,3-diene with Q2NHOAc 11

The reaction was carried out following general aziridination procedure A using 3-amino-2-isopropylquinazolin-4(3H)-one 11 (300 mg, 1.48 mmol), LTA (689 mg, 1.55 mmol), HMDS (597 mg, 3.70 mmol) and cyclohepta-1,3-diene (272 mg/0.30 cm³, 2.96 mmol) in dichloromethane (6 cm³). After work up the crude product was obtained as an off-white solid which was crystallised from ethyl acetate-light petroleum giving aziridine 32 (135 mg, 31%) as a white crystalline solid, mp 230–231 °C (Found: C, 72.9; H, 7.0; N, 14.2%. C₁₈H₂₂ON₃ requires C, 73.1; H, 7.1; N, 14.2%); $v_{\text{max}}/\text{cm}^{-1}$ 1660s and 1585m; δ_{H} 1.40 (3H, d, J 6.4, CH₃CHCH₃), 1.42 (3H, d, J 6.4, CH₃CHCH₃), 1.63–1.84 $(2H, struct. m, 2 \times CH), 2.04-2.19 (1H, m, 2 \times CH), 2.27-2.42$ (1H, m, CH), 2.51–2.64 (1H, m, CH), 2.94 (1H, br dd, J 8.0, ~5, azir. CH), 3.18 (1H, ddd, J 8.0, ~4, ~4, azir. CH), 3.65 [1H, heptet, J 6.4, CH(CH₃)₂], 5.95 (1H, ddd, J 11.4, 6.6, 3.2, CH=CH), 6.13 (1H, br ddd, J 11.4, ~5, ~2.5, CH=CH), 7.39 [1H, ddd, J 8.2, 6.8, 1.0, H-6(Q)], 7.62 [1H, dd, J 8.2, 1.0, H-8(Q)], 7.67 [1H, ddd, J 8.2, 6.8, 1.2, H-7(Q)] and 8.18 [1H, dd, J 8.2, 1.2, H-5(Q)]; δ_C 21.4, 21.5 [CH(CH_3)₂], 23.7, 28.9 $(2 \times \text{CH}_2)$, 31.1 [CH(CH₃)₂], 31.9 (CH₂), 50.6, 54.7 (2 × C-N), 121.7 [CCO(Q)], 122.9, 126.4, 126.5, 127.3, 133.8, 138.1 $[4 \times CH(Q), HC=CH], 146.6 [CN=C(Q)]$ and 160.5, 161.6 $[CN(Q), CO(Q)]; m/z(\%) 296 (MH^+, 100).$

Aziridination of cyclohepta-1,3-diene with Q¹NHOAc 6-TTB

General aziridination procedure **B** was followed in this reaction using Q¹NH₂ **5** (200 mg, 0.81 mmol), LTA (396 mg, 0.89 mmol), TTB (576 mg, 1.69 mmol) and cyclohepta-1,3-diene (152 mg/ 0.18 cm³, 2.96 mmol) in dichloromethane (5 cm³). Work up in the normal way gave a light green solid which was crystallised from ethyl acetate–light petroleum giving *aziridine* **33** (79 mg, 29%) as a colourless solid, mp 173–174 °C. [α]_D = +126.9 (c = 1.3, EtOH) (Found: MH⁺ 340.2025. C₂₀H₂₆O₂N₃ requires *M*H⁺ 340.2026); ν _{max}/cm⁻¹ 3490m, 1720s and 1580m; δ _H 1.02 [9H, s, C(CH₃)₃], 1.69–2.09 (4H, struct. m, 4 × CH), 2.30–2.45 (1H, m, CH), 2.77–2.88 (1H, m, CH), 2.90 (1H, dd, J 8.0, 4.8, azir. CH), 3.37 (1H, br ddd, J ~8, ~5, ~4, azir. CH), 3.77 (1H, d, J 10.4, CHOH), 5.08 (1H, d, J 10.4, CHOH), 6.01 (1H, ddd, J 11.9, 6.2, 2.0, C=CHCH₂), 6.10 (1H, ddd, J 11.9, 4.8, 2.5, NCHCH=C), 7.45 [1H, ddd, J 8.0, 6.9, 1.0, H-6(Q)], 7.64 [1H,

dd, J 8.2, 1.0, H-8(Q)], 7.71 [1H, ddd, J 8.2, 6.9, 1.0, H-7(Q)] and 8.21 [1H, dd, J 8.0 1.0, H-5(Q)]; $\delta_{\rm C}$ 23.1 (CH₂), 26.0 [C(CH₃)₃], 28.8, 31.7 (2 × CH₂), 38.4 [C(CH₃)₃], 52.4, 54.4 (2 × C-N), 74.5 (COH), 121.5 [CCO(Q)], 121.6, 126.3, 126.7, 126.9, 133.8 [4 × CH(Q), HC=CH], 139.5 (HC=CH), 144.7 [CN=C(Q)] and 157.6, 159.5 [CN(Q), CO(Q)]; m/z(%) 340 (MH⁺, 100), 307 (78), 289 (39) and 215 (32). A crystal suitable for X-ray structure determination was obtained from methanol.

Aziridination of cycloheptatriene with Q2NHOAc 11

General aziridination procedure A was followed in this reaction using 3-amino-2-isopropylquinazolin-4(3H)-one 11 (300 mg, 1.48 mmol), LTA (687 mg, 1.55 mmol), HMDS (597 mg, 3.69 mmol) and cycloheptatriene (272 mg, 0.29 cm³, 2.96 mmol) in dichloromethane (6 cm³) to give the crude product as a yellow oil. Column chromatography (4:1 light petroleum-ethyl acetate) gave aziridine 34 (193 mg, 45%), R_f 0.42, as a colourless oil which crystallised from ethyl acetate-light petroleum, mp 89-90 °C (Found: MH⁺ 294.1606. C₁₈H₂₀ON₃ requires MH⁺ 294.1606); $v_{\text{max}}/\text{cm}^{-1}$ 1660s and 1590s; δ_{H} 1.42 [6H, d, J 6.6, (CH₃)₂], 2.84 (1H, m, CHH), 2.99 (1H, dd, J 8.0, 3.7, azir. CHC=C), 3.03 (1H, ddd, J 15.6, ~5, ~5, CHH), 3.34 (1H, ddd, $J \sim 8, \sim 7, 5.0$, azir. CHCH₂), 3.63 [1H, heptet, J 6.6, CH(CH₃)₂], 5.89-6.08 (3H, struct. m, $3 \times C=CH$), 6.47 (1H, dd, J 11.0, 3.7, C=CH), 7.39 [1H, ddd, J 8.2, 6.6, 1.2, H-6(Q)], 7.58–7.68 [2H, m, H-7, H-8(Q)] and 8.17 [1H, dd, J 8.2, 1.0, H-5(Q)]; δ_C 21.1 $[CH(CH_3)_2]$, 28.7 (CH_2) , 30.8 $[CH(CH_3)_2]$, 48.9, 58.3 $(2 \times C-N)$, 121.2 [CCO(O)], 126.1, 127.0, 127.7, 127.8, 130.6, 130.9 $[3 \times CH(Q), 2 \times HC = CH], 133.5 [CH(Q)], 146.1 [CN = C(Q)]$ and 160.1, 161.1 [CN(Q), CO(Q)]; m/z(%) 294 (MH+, 100), 189 (39) and 173 (31).

Aziridination of cyclohexa-1,3-diene with Q²NHOAc 11 in acetonitrile

A modification of general aziridination procedure A was followed using 3-amino-2-isopropylquinazolin-4(3H)-one (200 mg, 0.98 mmol), LTA (458 mg, 1.04 mmol), HMDS (397 mg, 2.46 mmol) and cyclohexa-1,3-diene (158 mg/0.18 cm³, 1.97 mmol) in acetonitrile (4 cm³). After work up the crude product was obtained as a yellow oil. Column chromatography (2:1 light petroleum-ethyl acetate) gave alcohol 30 (74 mg, 27%) as a colourless oil, R_f 0.88 (Found: MH⁺ 300.1712. $C_{17}H_{22}O_2N_3$ requires MH^+ 300.1712); $v_{\text{max}}/\text{cm}^{-1}$ 3420m, 3300m, 1660s, 1615m and 1590s; $\delta_{\rm H}$ 1.35 (3H, d, J 6.9, CH₃CHCH₃), 1.41 (3H, d, J 6.9, CH₃CHCH₃), 1.85 (1H, struct. m, CH), 2.05-2.37 (3H, m, CH_2CHH), 2.85 (1H, dddd, $J \sim 10$, ~ 10 , ~ 3 , ~ 3 , CHNH), 3.64 [1H, heptet, J 6.9, $CH(CH_3)_2$], 3.96 (1H, br s, CHOH), 4.83 (1H, br s, OH), 5.78-5.87 (2H, struct. m, CH=CH), 5.93 (1H, d, J10, NH), 7.46 [1H, ddd, J8.0, 6.6, 1.3, H-6(Q)], 7.69-7.76 [2H, m, H-7, H-8(Q)] and 8.23 [1H, dd, J 8.0, 1.0, H-5(Q)]; $\delta_{\rm C}({\rm CDCl_3}, 62.9 {\rm MHz})$ 20.5 (CH₃CH*C*H₃), 21.6 (CH₂), 21.8 (CH₃CHCH₃), 26.0 (CH₂), 30.8 [CH(CH₃)₂], 45.0 (C-N), 62.6 (C-OH), 119.8 [CCO(Q)], 126.5, 126.6, 127.4, 131.3, 134.5 $[4 \times CH(Q), HC=CH], 147.2 [CN=C(Q)]$ and 162.5, 163.4 [CN(Q), CO(Q)]; *m/z*(%) 300 (MH⁺, 100), 282 (39), 204 (22), 189 (75) and 188 (50).

Further elution gave aziridine **20** (43 mg, 15%), R_f 0.57, as a colourless oil identical with that isolated previously and 2-isopropylquinazolin-4(3H)-one **31** (3 mg, 2%), R_f 0.22, as a colourless solid identical with an authentic sample.¹¹

Aziridination of cyclohexa-1,3-diene with Q¹NHOAc 6-TTB in acetonitrile

A modification of general aziridination procedure **B** was followed using Q¹NH₂ **5** (500 mg, 2.01 mmol), LTA (942 mg, 2.12 mmol), TTB (1.42 g, 4.20 mmol) and cyclohexa-1,3-diene (320 mg/0.38 cm³, 4.00 mmol) in acetonitrile (11 cm³). After work up, column chromatography (6:1 light petroleum–ethyl acetate) gave dienylamine **24** (19 mg, 7%) as a colourless oil, $R_{\rm f}$

0.26. Further elution gave aziridine 23 (101 mg, 38%) as a single diastereoisomer, R_f 0.21. The third fraction eluted, R_f 0.16, was a mixture which was re-chromatographed using Kieselgel (6:1 light petroleum-ethyl acetate) to give the major (Q^{l}) -amino alcohol diastereoisomer 27 (33 mg, 12%) as a colourless oil, $R_{\rm f}$ 0.19. $[a]_D$ +91.0 (c = 2.0, EtOH) (Found: MH⁺ 344.1975. $C_{19}H_{26}O_3N_3$ requires MH^+ 344.1975); v_{max}/cm^{-1} 3460s, 1660s, 1590s and 1470s; $\delta_{\rm H}$ (400 MHz) 1.05 [9H, s, (CH₃)₃], 1.65 (1H, dddd, J~13, ~7, ~3, ~3, CHHCN), 1.76 (1H, s, OH), 1.84 (1H, dddd, J 12.6, 11.9, 10.8, 5.0, CHHCN), 2.04 (1H, m, C=CCHH), 2.24 (1H, m, C=CCHH), 3.18 (1H, dddd, $J \sim 12$, ~8, ~3, ~3, CHNH), 3.62 (1H, br s, OH), 4.17 [1H, m, CH-(OH)CN], 5.16 (1H, br d, $J \sim 14$, CHOH), 5.75–5.86 (2H, m, CH=CH), 5.90 (1H, d, J7.6, NH), 7.50 [1H, ddd, J8.2, 7.1, 1.2, H-6(Q)], 7.62 [1H, dd, J 8.2, 1.2, H-8(Q)], 7.72 [1H, ddd, J 8.2, 7.1, 1.0, H-7(Q)] and 8.27 [1H, dd, J 8.2, 1.0, H-5(Q)]; m/z(%)344 (MH⁺, 100) and 233 (42). Irradiation at δ 4.17 resulted in loss of $J \sim 3$ Hz from the signal at δ 3.18 and simplification of the CH=CH multiplet.

Further elution gave the minor (Q^{l}) -amino alcohol diastereoisomer **28** (17 mg, 6%) as a colourless oil, R_f 0.14. $[a]_D$ +113.3 (c = 1.2, EtOH) (Found: MH⁺ 344.1975. $C_{19}H_{26}O_3N_3$ requires MH^+ 344.1975); $v_{\text{max}}/\text{cm}^{-1}$ 3420m, 1660s, 1595s and 1470m; $\delta_{\rm H}(400~{\rm MHz})~1.05~[9{\rm H},~{\rm s},~({\rm CH_3})_3],~1.80~(1{\rm H},~{\rm dddd},~J\sim12.5,\sim10,$ ~3, ~3, CHHCN), 1.92 (1H, dddd, J 12.5, 12.5, 8.5, 6.8, CHHCN), 2.19 (1H, struct. m, CHH), 2.31 (1H, struct. m, CHH), 2.96 (1H, dddd, J~13, ~10, ~3, ~3, CHNH), 3.60 (1H, d, J 10.5, Bu^tCHOH), 3.93 [1H, br dd, $J \sim 4$, ~ 3 , CH(OH)CN], 4.63 [1H, br d, J ~4, CH(OH)CN], 5.03 (1H, d, J 10.5, Bu^tCHOH), 5.79 (1H, dddd, J 9.9, ~5, ~1.5, ~1.5, CH=CCH₂), 5.84 (1H, d, J 10.0, NH), 5.90 (1H, ddd, J 9.9, ~5, ~3, C=CHCH₂), 7.55 [1H, ddd, J 8.2, 7.1, 1.2, H-6(Q)], 7.74 [1H, dd, J 8.0, 1.2, H-8(Q)], 7.83 [1H, ddd, J 8.0, 7.1, 1.2, H-7(Q)] and 8.30 [1H, dd, J 8.2, 1.2, H-5(Q)]; $\delta_{\rm C}$ (75 MHz) 22.2, 26.2 $(2 \times CH_2)$, 26.3 [(CH₃)₃], 38.5 (CHNH), 62.6 (COH), 75.0 (CHOH), 120.5 [CCO(Q)], 126.5, 127.2, 127.6, 127.7, 131.6, 135.4 [4 \times CH(Q), HC=CH], 146.3 [CN=C(Q)] and 158.8, 163.5 [CO(Q), CN(Q)]; m/z(%) 344 (MH⁺, 100) and 233 (30).

Irradiation of the signal at δ 3.93 converted that at δ 4.63 into a singlet and that at δ 5.79 into a ddd $J \sim 10$, ~ 1.5 , ~ 1.5 Hz, and that at δ 2.96 into a ddd J 13, 10 and 3.2 . Irradiation of the signal at δ 2.96 affected only that at δ 3.93 and CH_2 signals at δ 1.80 and 1.92.

Ring-opening of aziridine 23 with CH₃CN-H₂O-acid

Aziridine 23 (220 mg, 0.68 mmol) was stirred in acetonitrile (4 cm³) containing water (1 cm³) and toluene-p-sulfonic acid (8 mg) at ambient temperature for 2 h. Ethyl acetate (10 cm³) was added, the solution washed with saturated sodium hydrogencarbonate solution (10 cm³) and the organic layer separated, dried and evaporated. Column chromatography (7:1 light petroleum-ethyl acetate) of the residue gave cyclic ether 29 (53 mg, 24%) as a colourless oil, R_f 0.38. $[a]_D$ + 151.5 (c = 1.0, EtOH) (Found: MH⁺ 326.1869. $C_{19}H_{24}O_2N_3$ requires MH⁺ 326.1868); $v_{\text{max}}/\text{cm}^{-1}$ 1660s, 1600s, 1520m and 1470s; δ_{H} 1.30 [9H, s, $(CH_3)_3$], 1.92–2.37 (4H, m, 2 × CH_2), 3.23 (1H, struct. m, CHNH), 4.53 (1H, struct. m, HCO), 4.81 (1H, s, ButCHO), 5.78 (1H, m, incl. $J \sim 10$, CH=CH), 6.06 (1H, struct. m, incl. $J \sim 10$, CH=CH), 6.60 (1H, br s, NH), 7.48 [1H, ddd, J 8.2, 5.3, 3.2, H-6(Q)], 7.69-7.76 [2H, m, H-7 and H-8(Q)] and 8.25 [1H, ddd, J 8.2, 1.2, H-5(Q)]; $\delta_{\rm C}$ 20.8, 26.3 (2 × CH₂), 27.2 [(CH₃)₃], 34.9 $[C(CH_3)_3]$, 56.2 (C-N), 74.4, 76.9 (2 × CHO), 120.4 [CCO(Q)], 126.6, 126.9, 127.2, 128.6, 132.4, 134.2 [4 × CH(Q), HC=CH], 146.7 [CN=C(Q)] and 154.1, 160.6 [CN(Q), CO(Q)]; m/z(%) 326 (MH⁺, 100), 246 (30) and 215 (48).

Further elution gave the major alcohol diastereoisomer 27 (114 mg, 49%) R_f 0.27, and the minor diastereoisomer **28** (31 mg, 13%) $R_{\rm f}$ 0.14, both as colourless oils identical with those isolated previously.

Ring-opening of aziridine 23 in acetonitrile containing H₂¹⁸O

Aziridine 23 (dr 5:1) (50 mg, 0.15 mmol) was dissolved in acetonitrile (1 cm³) and $H_2^{18}O$ (8 μ l, 2.5 equiv.) was added followed by toluene-p-sulfonic acid (~2 mg). After stirring for 45 min, ethyl acetate (10 cm³) was added, the solution washed with saturated aqueous sodium hydrogencarbonate (10 cm³) and the organic layer separated, dried, and evaporated. An NMR spectrum of the residue showed the presence of unchanged aziridine 23 (δ 3.58) and (Q¹)-amino alcohol 27 and **28** (δ 3.18 and 2.95; ratio 2:1 respectively) in a ratio of 4:1. For this mixture, mass spectrometry showed no ¹⁸O incorporation into the aziridine (MH⁺ 326). For the mixture of (Q¹)amino alcohols 27 and 28 (Found: MH⁺ 346.2016. C₁₉H₂₆- $N_3^{16}O^{18}O$ requires MH^+ 346.2017) the ratio of 344:346 was 2:1. Separation of unchanged aziridine 23 and (Q1)-amino alcohols 27 and 28 using flash silica [base-washed with triethylamine (2%) solution in light petroleum-ethyl acetate and mass spectroscopy on the recovered alcohol 28 showed the ¹⁶O: ¹⁸O ratio was raised to 8:1. The recovered aziridine 23 (dr 14:1) was redissolved in acetonitrile (1 cm³) and stirred for a further 2 h with $H_2^{18}O$ (5 µl, 2.5 equiv.) and toluene-p-sulfonic acid (~2 mg). After work up as described above, NMR spectroscopy showed the presence of alcohol 27 (Found: MH+ 346.2017. $C_{19}H_{26}N_3^{16}O^{\bar{1}8}O$ requires MH^+ 346.2017) with a 344 : 346 ratio of 1:1 (10:1 after purification by flash chromatography).

Conversion of (Q²)-amino alcohol 30 into oxazolidinone 36

(Q²)-amino alcohol 30 (60 mg, 0.20 mmol) was heated under reflux in THF (1 cm³) with sodium hydride (5 mg, 0.22 mmol) and 1,1'-carbonyldiimidazole (45 mg, 0.30 mmol) under nitrogen for 3 h. After cooling, ethyl acetate (10 cm³) was added and the solution washed with saturated sodium hydrogencarbonate solution (10 cm³), the organic layer was separated, dried and evaporated to give the crude product as a colourless solid. Column chromatography (5:1 light petroleum-ethyl acetate) gave unchanged alcohol 30 (21 mg, 35%) as a colourless oil, $R_{\rm f}$ 0.49.

Further elution gave oxazolidinone 36 (35 mg, 53%) as a colourless solid (R_f 0.33), mp 148-149 °C (from ethanol) (Found: MH⁺ 326.1505. $C_{18}H_{20}O_3N_3$ requires MH⁺ 326.1504); $v_{\rm max}/{\rm cm}^{-1}$ 1770s, 1680s and 1600s; $\delta_{\rm H}$ 1.34 (3H, d, J 6.6, CH₃CHCH₃), 1.40 (3H, d, J 6.6, CH₃CHCH₃), 1.86 (2H, m, CHH), 2.08-2.21 (1H, m, CHH), 2.24-2.40 (1H, m, CHH), 3.18 [1H, heptet, J 6.6, CH(CH₃)₃], 4.02–4.15 (1H, m, NCH), 5.00–5.07 (1H, m, HCO), 6.00 (1H, m, incl. J 10.2, CH=CH), 6.26-6.35 (1H, m, incl. J 10.2, CH=CH), 7.45 [1H, ddd, J 8.0, 6.8, 1.4, H-6(Q)], 7.70 [1H, dd, J 8.2, 1.4, H-8(Q)], 7.77 [1H, ddd, J 8.2, 6.8, 1.1, H-7(Q)] and 8.22 [1H, dd, J 8.0, 1.1, H-5(Q)]; $\delta_{\rm C}$ 20.2 (CH₂), 21.0 (CH₃), 22.8 (CH₂), 22.9 (CH₃), 30.6 [CH(CH₃)₃], 54.4 (NCHCH₂), 71.1 (C=CHCO), 121.4 [CCO(Q)], 123.2, 127.1, 127.3, 127.9, 133.3, 135.4 [4 × CH(Q), HC=CH], 147.4 [CN=C(Q)], 157.6, 159.9 and 163.4 [C=O, CN(Q), CO(Q)]; m/z(%) 326 (MH⁺, 66), 307 (100) and 289 (50).

Conversion of (Q¹)-amino alcohol 27 into oxazolidinone 37

(Q1)-amino alcohol 27 (37 mg, 0.11 mmol) was heated under reflux in THF (1 cm³) with sodium hydride (3 mg, 0.12 mmol) and 1,1'-carbonyldiimidazole (22 mg, 0.14 mmol) under nitrogen for 3 h. After cooling, ethyl acetate (10 cm³) was added and the solution washed with saturated sodium hydrogencarbonate solution (10 cm³), the organic layer was separated, dried and evaporated to give the crude product as a colourless solid. Column chromatography (5:1 light petroleum-ethyl acetate) gave unchanged alcohol 27 (16 mg, 43%) as a colourless oil, R_f 0.18.

Further elution gave oxazolidinone 37 (20 mg, 49%) as a colourless oil, $R_f 0.10$. $[a]_D + 39.5$ (c = 1.0, EtOH) (Found: MH⁺ 370.1766. $C_{20}H_{24}O_4N_3$ requires MH^+ 370.1767); v_{max}/cm^{-1} 1790m, 1705m and 1605m; $\delta_{\rm H}$ 0.99 [9H, s, (CH₃)₃], 1.67–1.86 (2H, m, CH_2), 2.00–2.24 (2H, m, CH_2), 3.32 (1H, d, J 10.8, CHOH), 4.12 (1H, ddd, J 11.6, 6.7, 4.6, NCH), 4.43 (1H, d, J 10.8, CHOH), 4.89–4.95 (1H, m, HCO), 5.93 (1H, dddd, J ~10, ~3, ~3, ~3, CH=CH), 6.18–6.27 (1H, m, CH=CH), 7.45 [1H, ddd, J 8.0, 6.9, 1.0, H-6(Q)], 7.62 [1H, dd, J 8.0, 1.0, H-8(Q)], 7.73 [1H, ddd, J 8.0, 6.9, 1.0, H-7(Q)] and 8.21 [1H, dd, J 8.0, 1.0, H-5(Q)]; m/z(%) 370 (MH $^+$, 100), 344 (22), 307 (25), 215 (28).

Conversion of (O1)-amino alcohol 28 into oxazolidinone 38

(Q¹)-amino alcohol **28** (23 mg, 67 μmol) was heated under reflux in THF (1 cm³) with sodium hydride (2 mg, 73 μmol) and 1,1'-carbonyldiimidazole (16 mg, 0.10 mmol) under nitrogen for 3 h. After work up as described above, column chromatography (3:1 light petroleum–ethyl acetate) of the crude product gave alcohol **28** (11 mg, 48%) as a colourless oil, $R_{\rm f}$ 0.24.

Further elution gave oxazolidinone 38 (11 mg, 44%) as a colourless oil, R_f 0.19 (Found: MH⁺ 370.1766. C₂₀H₂₄O₄N₃ requires MH^+ 370.1767); $v_{\text{max}}/\text{cm}^{-1}$ 1790 m, 1695 m and 1605 m; $\delta_{\rm H}$ 1.06 [9H, s, (CH₃)₃], 1.85–2.00 (2H, m, 2 × CH), 2.10– 2.27 (1H, m, CHH), 2.37-2.54 (1H, m, CHH), 2.95 (1H, d, J 11.2, CHOH), 4.54 (1H, ddd, J~8, ~4, ~4, NCH), 4.71 (1H, d, J 11.2, CHOH), 5.23-5.31 (1H, m, HCO), 5.89 (1H, dddd, J 13.1, ~3, ~3, ~2, CH=C), 6.18–6.27 (1H, m, C=CH), 7.52 [1H, ddd, J 8.2, 6.8, 1.1, H-6(Q)], 7.70 [1H, dd, J 8.2, 1.1, H-8(Q)], 7.82 [1H, ddd, J 8.2, 6.8, 1.1, H-7(Q)] and 8.27 [1H, dd, J 8.2, 1.1, H-5(Q)]; $\delta_{\rm C}$ (75 MHz) 20.3, 21.9 (2 × CH₂), 26.3 [(CH₃)₃], 37.2 $[C(CH_3)_3]$, 70.9, 74.7 $(2 \times CH)$, 121.7 [CCO(Q)], 122.8, 127.7, 127.9, 128.0, 133.2, 135.7 [$4 \times CH(Q)$, HC=CH], 146.2 [CN=C(Q)], 157.2 [OC(O)N] and 159.0, 159.6 [CN(Q), CO(Q)]; m/z(%) 370 (MH⁺, 100). A COSY spectrum showed the signal at δ 1.85–2.00 (CH₂) coupled with δ 4.54 (NCH) and δ 5.89 $(CH=CCH_2)$ coupled with δ 5.23–5.31 (HCO).

Q¹-N reductive cleavage of oxazolidinone 37 with Sm(II) iodide ¹⁰

A two-necked round bottom flask fitted with a 3-way tap and a septum cap was flame dried, flushed with argon and oxazolidinone 37 (17 mg, 45.9 μmol) dissolved in THF (1 cm³) was added via a syringe followed by tert-butyl alcohol (0.5 cm³) in THF (0.5 cm³). Samarium(II) iodide (0.1 M solution in THF) was then added dropwise until the dark blue colour of samarium(II) persisted (~1 cm³). Ethyl acetate (5 cm³) was added and the solution washed with saturated sodium hydrogencarbonate solution (5 cm³), the organic layer was separated, dried and evaporated to give the crude product as a yellow semi-solid. Column chromatography (6:1 light petroleum–ethyl acetate) gave a mixture of products. Subsequent Kieselgel chromatography (9:1 ethyl acetate–methanol) gave quinazolin-4(3H)-one 40 (8 mg, 75%) as a colourless crystalline solid, identical with a sample isolated previously.⁵

Further elution gave *oxazolidinone* **39** (4 mg, 62%) as a colourless solid, mp 85–86 °C (from light petroleum–ethyl acetate) [lit. ¹³ (racemate) 86–88 °C]. [a]_D -10.6 (c = 0.5, EtOH) (Found: MH $^+$ 140.0711. C₇H₉O₂N requires MH $^+$ 140.0712); $\nu_{\rm max}/{\rm cm}^{-1}$ 3220m and 1740m; $\delta_{\rm H}$ 1.70–1.78 (1H, m, CH), 1.80–2.05 (2H, m, 2 × CH), 2.15–2.31 (1H, m, CH), 3.89–4.03 (1H, struct. m, CHNH), 4.87–4.95 (1H, m, HCO), 5.07 (1H, br s, NH), 5.79–5.88 (1H, m, CH=CH) and 6.11–6.22 (1H, m, CH=CH); m/z(%) 162 (MNa $^+$, 100) and 140 (MH $^+$, 67).

Q1-N reductive cleavage of oxazolidinone 38 with Sm(II) iodide 10

The procedure described above was carried out using oxazolidinone **38** (11 mg, 29.7 μmol), and *tert*-butyl alcohol (0.5 cm³) in THF (1 cm³). Samarium(II) iodide (0.1 M solution in THF) was added dropwise until the dark blue colour of samarium(II) metal persisted (~0.8 cm³). After work up column chromatography (6:1 light petroleum–ethyl acetate) of the yellow

residue gave a mixture of products. Subsequent Kieselgel chromatography (9:1 ethyl acetate–methanol) gave quinazolin-4(3H)-one **40** (5 mg, 72%), $R_{\rm f}$ 0.71, as a colourless crystalline solid identical to that isolated previously.

Further elution gave oxazolidinone **enant. 39** (2.5 mg, 60%) as a colourless solid, mp 87–88 °C (from light petroleum–ethyl acetate). [a]_D +13.2 (c = 0.5, EtOH) (Found: MH $^+$ 140.0711. C₇H₉O₂N requires MH $^+$ 140.0712); v_{max}/cm^{-1} 3220m and 1740m; NMR spectrum identical to that of **39** above; m/z(%) 162 (MNa $^+$, 100) and 140 (MH $^+$, 67).

Crystal structure determinations of 21 and 33

Data for both compounds were measured on a Siemens P4 diffractometer with graphite monochromated Mo-K α radiation ($\lambda=0.7107\,$ Å) using an ω -scan technique. Three standard reflections monitored every 100 scans showed no significant variation in intensity; the reflections were corrected for Lorentz and polarisation effects.

The structures were solved by direct methods and refined by full-matrix least squares on F^2 using the program SHELXL-97. All hydrogen atoms were included in calculated positions (C–H = 0.96 Å) using a riding model. All non-hydrogen atoms were refined as anisotropic. The absolute configuration of 33 at C1 is determined by the preparation from the (S)-tert-leucine.

Crystal data for 21. $C_{15}H_{12}F_3N_3O$, M=307.28, monoclinic, space group C2/c, a=24.012(3), b=7.609(1), c=15.758(2) Å, $\beta=105.89(1)^\circ$, V=2769.1(6) ų, T=200 K, Z=8, $\mu(\text{Mo-K}\alpha)=0.123$ mm $^{-1}$, colourless block, crystal dimensions $0.49\times0.21\times0.20$ mm. Full matrix least squares based on F^2 gave R1=0.070 for 808 observed data $(F>4\sigma(F))$ and wR2=0.232 for all 1038 data, GOF=1.078 for 145 parameters.

Crystal data for 33. $C_{20}H_{25}N_3O_2$, M = 339.43, tetragonal, space group $P4_3$, a = b = 10.242(2), c = 17.385(3) Å, V = 1823.7(6) Å³, T = 200 K, Z = 4, $\mu(\text{Mo-K}\alpha) = 0.08$ mm⁻¹, colourless block, crystal dimensions $0.62 \times 0.48 \times 0.41$ mm. Full matrix least squares based on F² gave R1 = 0.036 for 1579 observed data $(F > 4\sigma(F))$ and wR2 = 0.097 for all 1779 data, GOF = 1.030 for 226 parameters.

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