

Fig. 2 Target library of cysteine-derived tetramates.

spectroscopic (XPS) analysis of some derivatives clearly identified the presence of Ca(II)<sup>29</sup> in metal-chelated tetramates and a high abundance of Ca(II), together with Na(I), Mg(II), Fe(III) and Zn(II), was also found using inductively coupled plasma mass spectrometry (ICP-MS).<sup>30</sup> The metal chelating properties of tetramates are most immediately evident by post-chromatographic signal broadening in <sup>1</sup>H NMR spectra, which can only be removed by acid wash,<sup>29–32</sup> and the metal complexation behaviour of tetramates was also found to be both structure and metal-ion dependent.<sup>33</sup> We have described earlier that cysteine-derived bicyclic tetramates **1** are excellent metal chelating agents, and established that the C-6, C-8 and/or C-9 carbonyl groups were directly involved in that metal complexation, but that it could be blocked by protection of the enolic system;<sup>34</sup> it seemed that modification of the C-5 ester (CO<sub>2</sub>R<sup>1</sup>, Fig. 1), along with the concave-convex shape of the bicyclic tetramate system, might be further exploited to alter the metal binding ability of the tetramate system by steric interference distal to the region of metal chelation.<sup>33</sup>

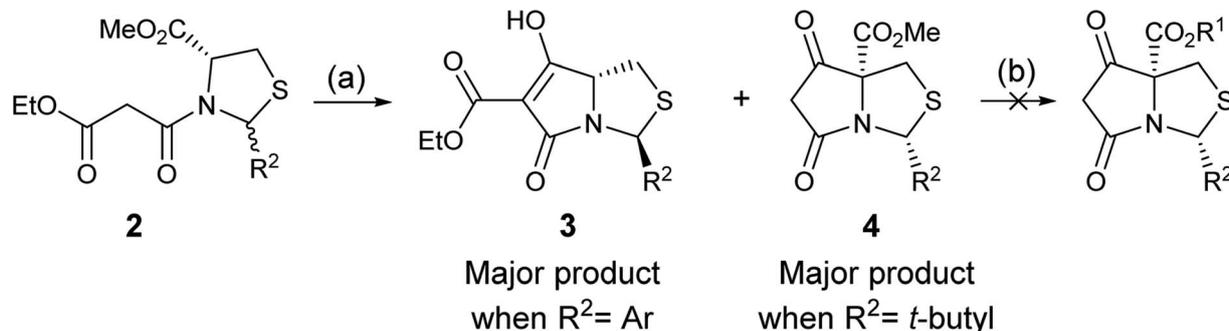
Specifically, target compounds were designed with increasing steric bulk at C-5, with R<sup>1</sup> varying from methyl to allyl, isopropyl, cyclohexyl, benzyl, nitro-benzyl and bromo-benzyl esters (Fig. 2). Polar pyridyl rings R<sup>2</sup> were incorporated at C-2 to increase overall hydrophilicity. Various C-9 side chains R<sup>3</sup> such as cyclohexyl, phenyl, phenethyl, styrenyl, and heterocyclic pyridyl and thiazolyl systems in tetramate ketones as well as cyclohexyl and adamantyl amide pendants in tetramate carboxamides were examined (Fig. 2). The cyclohexyl ketone pendant is a simplified mimic of the *trans*-decalin system found in many tetramate-containing natural products, such as kibdelomycin, altersetin, zopfiellamide A and signermycin B. Phenethyl and styrenyl derivatives were also structurally similar to naturally occurring tetramates with alkyl ketone side chains

such as epicoccarine A, streptolydigin and virgineone, as well as the endogenous substrates of undecaprenyl pyrophosphate synthase (UPPS) such as isopentenyl pyrophosphate (IPP) and farnesyl pyrophosphate (FPP); UPPS is a known biological target of tetramates.<sup>35–37</sup> Thus, a C-9 ethyl ketone series and a C-9 heterocyclic ketone series was synthesised to compare structural activity relationships, to demonstrate the versatility of the synthetic route for functionalisation.

## Results and discussion

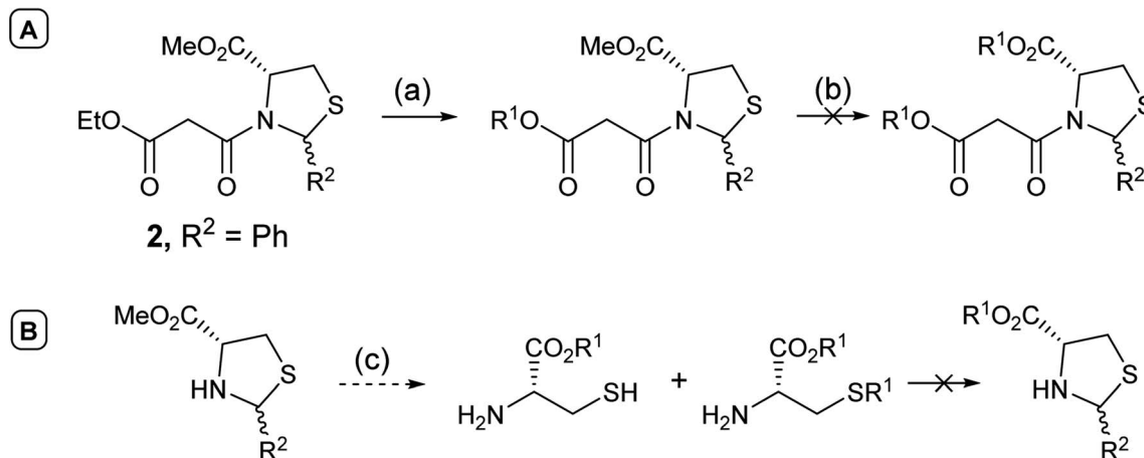
In order to access C-5 ester variants, our initial approach sought to exploit the fact that *N*-acylthiazolidines **2** with aromatic C-2 substituents (R<sup>2</sup> = Ar or HetAr) selectively cyclise from the side chain enolate onto the C-5 methyl ester to form tetramate esters **3** as the major product, leaving no C-5 substituents, but that *N*-acylthiazolidines (R<sup>2</sup> = *t*-Bu) with a bulkier C-2 *t*-butyl group cyclise from the enolate of the C-5 methyl ester to form the tetramate core **4**; unfortunately, however, the methyl ester of the latter could not be directly transesterified, suggesting unusual stability, probably arising from the significant steric hindrance of the bicyclic system (Scheme 1).

Alternatively, attempts to adjust the identity of the ester earlier in the synthetic sequence, by for example, direct transesterification of *N*-acylthiazolidine **2** (R<sup>2</sup> = Ph), in fact favoured reaction at the less hindered ethyl ester on the side chain (Scheme 2A); transesterification of the C-5 ester group could not be achieved, with the harsh conditions resulting instead in ring opening and degradation of the *N*-acylthiazolidine **2** (R<sup>2</sup> = Ph). Transesterification of the unprotected thiazolidine, known to be less stable than the *N*-acylthiazolidine derivative, not surprisingly led to ring opening; only transesterified *L*-cysteine and the

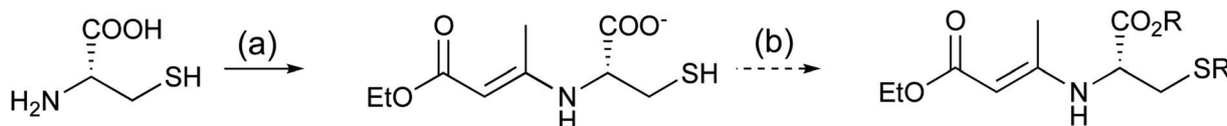


Scheme 1 Reagents and conditions: (a) KO<sup>t</sup>Bu, THF, reflux, 4 h; (b) R<sup>1</sup>OH, catalytic *p*-TsOH monohydrate or DMAP, toluene, reflux.





**Scheme 2** (A) Direct transesterification of *N*-acylthiazolidine; (B) transesterification of the unprotected thiazolidine. Reagents and conditions: (a)  $R^1OH$ ,  $R^1 = p$ -methoxybenzyl, catalytic DMAP, toluene, reflux, o.n.; (b) additional equiv. of  $R^1OH$ , toluene, reflux, o.n.; (c)  $R^1OH$ ,  $R^1 =$  benzyl or  $p$ -methoxybenzyl, catalytic  $p$ -TsOH monohydrate or DMAP, toluene, reflux, o.n.



**Scheme 3** Reagents and conditions: (a) *N,N,N,N*-tetramethylguanidine, anhydrous DMF, 0 °C for 30 min followed by addition of ethyl acetoacetate, 0 °C to r.t., 18 h; (b) RBr, (R = benzyl), r.t., 24 h.

product with alkylation on the thiol group were detected by mass spectrometry (Scheme 2B).

With the failure to directly functionalise both thiazolidines and the derived tetramate core at the C-5 ester group, it became apparent that access to esters of *L*-cysteine at the outset would be required. The formation of such systems is complicated by the presence of the highly nucleophilic thiol group, and while application of an elegant literature method to prepare esters of *L*-serine and *L*-threonine<sup>38</sup> using amine protection with ethyl acetoacetate followed by esterification with alkyl halides under basic conditions and subsequent amine deprotection with methanolic HCl indeed gave the desired ester, it was also obtained along with unwanted reaction at the thiol side chain (Scheme 3).

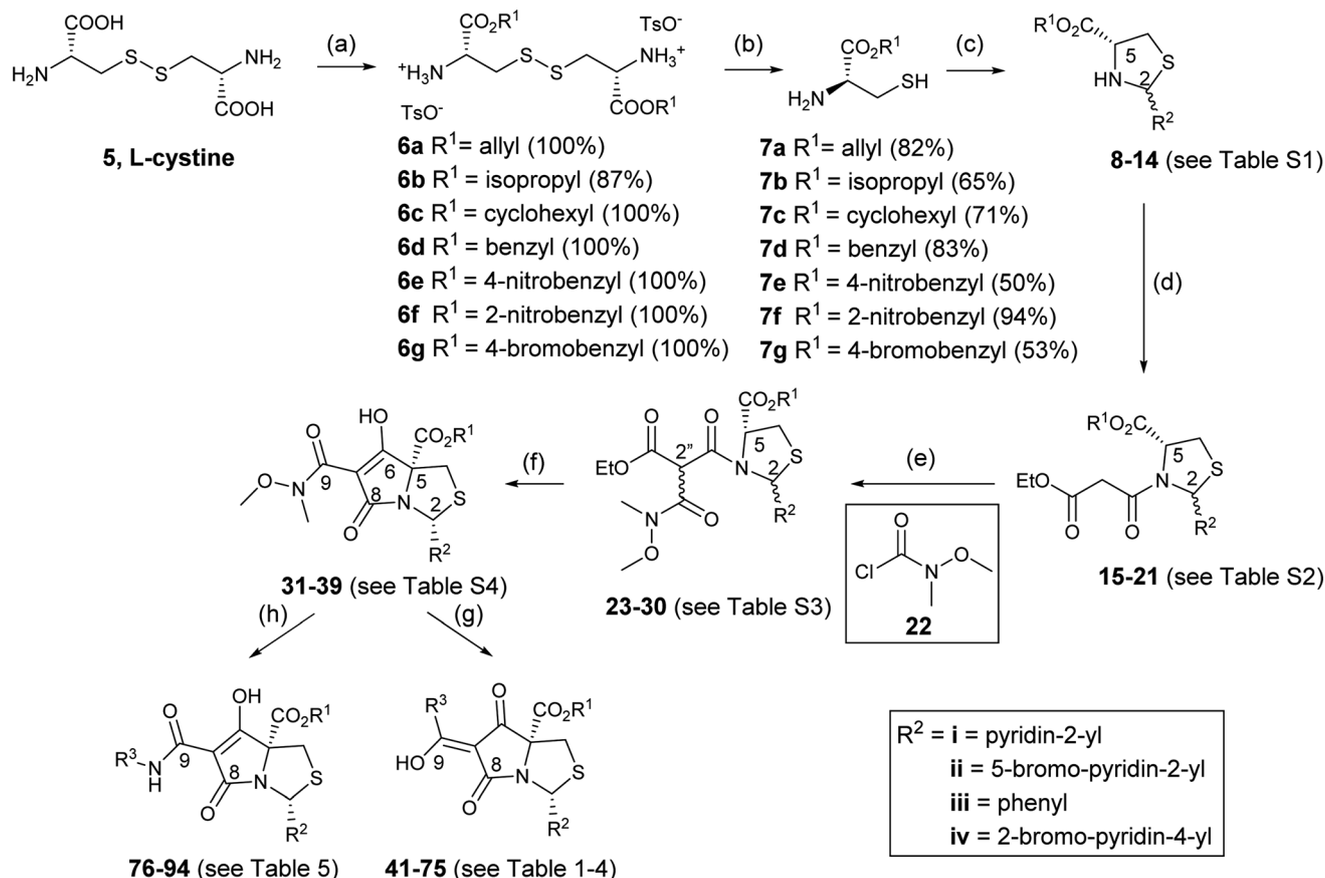
We therefore focussed on a strategy exploiting the attenuated nucleophilicity of sulphur in the disulphide group of *L*-cystine 5. Esterification of *L*-cystine by thionyl chloride and simple alcohols occurred in quantitative yield,<sup>39</sup> although reaction with bulkier alcohols such as benzyl alcohol and *p*-methoxybenzyl alcohol was more limited and gave incomplete reaction, with starting material mostly being recovered. Acid-catalysed esterification with *p*-TsOH monohydrate in toluene proved to be more reliable,<sup>40,41</sup> and was even better using cyclohexane as solvent,<sup>42–44</sup> giving diesters of *L*-cystine **6a–g** which precipitated as their bis(*p*-toluenesulphonate) salts (Scheme 4).<sup>45–47</sup> Although widely used as starting materials for synthesising biologically relevant molecules, these systems have generally been incompletely characterised.<sup>46–52</sup>

These diesters of *L*-cystine **6a–g** were then reduced at the disulphide bond to produce esters of *L*-cysteine **7a–g**. Reduction by sodium borohydride by the literature method<sup>53</sup> also gave unwanted reduction of the ester groups, but fully selective cleavage of the disulphide bond was successfully achieved with dithiothreitol (DTT) under weakly alkaline conditions ( $Et_3N$ ) (Scheme 4) in good yield.<sup>45,54–56</sup>

With these materials reliably in hand, condensation of *L*-cysteine esters **7a–g** with aromatic aldehydes ( $R^2CHO$ ) gave the corresponding thiazolidines **8–14** as a mixture of *cis/trans*-2,5 diastereomers (ratio varying from 0.5 : 1 to 4.5 : 1) in good yield (Scheme 4 and Table S1, ESI†). The characteristic and highly conserved H-2 and H-5 chemical shifts were consistent with those previously found in C-5 methyl ester thiazolidines,<sup>33</sup> both being more downfield in the *trans*-diastereomers.

Using DCC/DMAP coupling conditions, thiazolidines **8–14** were converted to the corresponding *N*-acylthiazolidines **15–21** as a mixture of *cis/trans*-2,5 diastereomers in high yield (Scheme 4 and Table S2, ESI†). In the NMR spectrum, complexity arose since each diastereomer appeared as a rotameric pair, and H-2 chemical shifts were consistent with that observed previously in *N*-acylthiazolidines with C-2 aromatic substituents.<sup>33</sup> Thus, in the C-2 phenyl series, the H-2 chemical shifts for the major *trans* rotamers were consistently more downfield; this trend was reversed in the minor rotamers. In the C-2 pyridyl series, the H-2 chemical shifts for major *trans* rotamers were consistently more upfield, and this trend was preserved in the minor rotamers. In the case of **17i**, two diastereomers of the starting thiazolidine



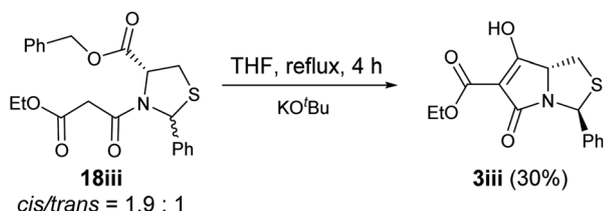


**Scheme 4** Reagents and conditions: (a) *p*-TsOH monohydrate,  $\text{R}^1\text{OH}$ , cyclohexane, reflux, 5–16 h; (b) DTT,  $\text{Et}_3\text{N}$ , DCM,  $\text{N}_2$ , r.t., 24 h; (c)  $\text{R}^2\text{CHO}$ , petrol 40/60, reflux, 18 h; (d) mono-ethyl malonate, DCC, DMAP, DCM, 0 °C to r.t., 18 h; (e) *N*-methoxy-*N*-methylcarbamoyl chloride **22**, dry pyridine,  $\text{MgBr}_2$ , DCM, 0 °C to r.t., 18–24 h; (f) DBU dry THF, r.t., 24 h; (g)  $\text{R}^3\text{MgBr}$  (prepared fresh with  $\text{R}^3\text{Br}$ , Mg, dibromoethane, dry THF, reflux, 1 h), dry THF, –15 °C, 1 h or o.n.; (h)  $\text{R}^3\text{NH}_2$ , THF/toluene, reflux, 18 h.

**10i** could be separated by column chromatography and each was taken for separate *N*-acylation. The reaction with *cis*-**10i** produced a mixture of predominantly *cis*-malonamide **17i** (*cis/trans* ratio of 4.8 : 1), while the reaction with *trans*-**10i** yielded a mixture of predominantly *trans*-malonamide **17i** (*cis/trans* ratio of 0.2 : 1). That these two separate reactions starting from a pure single diastereomer produced a mixture of diastereomers, and that the *cis/trans* ratio changes from thiazolidines **8–14** to *N*-acylthiazolidines **15–21**, indicated the existence of an equilibrium between the two thiazolidine diastereoisomers of **8–14** under the reaction conditions. It was generally observed that when groups at C-2 and C-5 became bulkier, as in the cases **18–21**, the production of the more stable *cis*-diastereomer was

favoured, presumably since in that case, both groups were able to assume pseudodiequatorial positions.

Noteworthy is that when *N*-acylthiazolidine **18iii** (with a bulkier C-5 benzyl ester) was subjected to Dieckmann cyclisation, it cyclised from the more stable side chain enolate to the C-5 ester, giving tetramate ester **3iii** as the major product and therefore with loss of the C-5 ester (Scheme 5); the cyclisation started with a 1.9 : 1 mixture of *cis/trans* diastereomers, and while the *trans* diastereomer of **18iii** cyclised to form **3iii**, the *cis* diastereomer was recovered partially (degradation was observed). This chemoselectivity in cyclisation has been consistently observed in *N*-acylthiazolidines with C-2 aryl substituents.<sup>32</sup> Since this reaction outcome was the undesired chemoselectivity, to avoid this, a Weinreb amide was introduced under Sucunza's acylation conditions,<sup>57</sup> an approach which had been used previously for the acylation of *N*-acyloxazolidines or thiazolidines with a C-2 *t*-butyl group.<sup>29</sup> Successful C-acylation on the side chain of the *N*-acylthiazolidines **15–21** was achieved to give **23–30** (Scheme 4 and Table S3, ESI†), but even after 24 hours, reactions were not complete, with starting malonamides enriched in *trans*-diastereomers being recovered, even with 1.5 equivalents of the acid chloride **22**. Each of products **23–30** was obtained as an approximately equal ratio



**Scheme 5**



mixture of C2'' *R/S* diastereomers (Table S3, ESI†). Consistent with previous observations,<sup>58</sup> the side chain tricarbonyl system existed in the keto-form, indicated by the presence of an H-2'' singlet in <sup>1</sup>H NMR spectra. Exclusive formation of the *cis* diastereomer was found in all cases except for 27ii. The relative stereochemistry was supported by NOE analysis (Fig. S1, ESI†), while 2D NOESY also confirmed the presence of rotamers in the mixture. In the case of 26i, the predominantly *cis* mixture of 17i (*cis/trans* = 4.8 : 1) produced *cis*-26i exclusively under the acylation conditions in 68% yield. The reaction with the predominantly *trans* mixture of 17i (*cis/trans* = 0.2 : 1) was less straightforward; thus, unreacted *trans* starting material 17i was recovered in 42% yield, and product 27a was formed in 38% yield as a mixture of *cis/trans* (0.4 : 1) and C2'' *R/S* (0.8 : 1) diastereomers. This result suggested that the *cis*-malonamides reacted more favourably under the acylation conditions and that there could be interconversion between *cis/trans* diastereomers to favour the *cis*-2,5 products. The diastereomers were not separable by flash column chromatography and the mixture was used directly for Dieckmann cyclisation (see below).

Using literature conditions with DBU,<sup>29</sup> successful Dieckmann cyclisation of 23–30 was achieved generally in good yield, leading to the desired tetramate Weinreb amides 31–39 with different C-5 esters (Scheme 4 and Table S4, ESI†). The presence of basic C-2 pyridyl groups (R<sup>2</sup>) and hence the corresponding increase in product polarity added to the difficulty of acidic work-up after the reactions, lowering yields significantly. For cyclisation with 24i and 25i, which contained a C-2 unsubstituted pyridyl ring, although the product was detected by mass spectrometry, no product could be extracted; increasing the lipophilicity of groups at C-2 and C-5 avoided this difficulty. Under these Dieckmann cyclisation conditions, the *cis* diastereomer of tricarbonyl malonamides cyclised from the C-5 enolate onto the side chain carbonyl by placing the C-2 substituents

at the less hindered *exo*-face of the bicyclic ring, producing tetramate derivatives 31–39 with retention of the C-5 ester groups. However, in the reaction with *trans*-27i, no tetramate product was observed and starting material was recovered, presumably due to the greater bulk of the C-5 ester. Noteworthy was the efficiency and chemoselectivity of the synthetic route, and with bulky C-5 esters, the *cis*-diastereomer was the only diastereomer that cyclised under the Dieckmann cyclisation, retaining the C-5 ester. The basic conditions with DBU also helpfully favoured exclusively reaction at the ester of the side chain, so that the Weinreb amide remained available for late-stage functionalisation at C-7. Weinreb amides 31–39 were completely enolic, with no C-7 proton observed in <sup>1</sup>H NMR spectra, and in the *endo*-enolic tautomeric pair **AB** form (Fig. 3 and Scheme 4), as supported by HMBC analysis (Fig. S2, ESI†) and by comparison of carbonyl <sup>13</sup>C NMR peaks with earlier reports.<sup>29</sup> They were also a single diastereomer at C-2/C-5 with characteristic and consistent H-2 chemical shifts (Table S4, ESI†). The *cis* relationship between C-5 ester and C-2 aromatic rings was supported by NOE analysis (Fig. S2, ESI†).

In the reactions of 28iv and 29iv, which contained only C-5 nitrobenzyl esters, the product was unexpectedly a 1 : 1 inseparable mixture of tetramate Weinreb amides with the respective C-5 nitrobenzyl ester along with ethyl ester 37iv (Table S4, ESI†). The latter by-product 37iv would arise from transesterification with ethoxide generated under the basic conditions of the reaction. Similar by-product formation was observed from the crude product of the reaction of 35 and 39 but to a much less extent and desired products could be easily purified with column chromatography. The by-product formation with bulky C-5 esters limited the scope of the process, and made 39 all the more valuable, since the bromo-substituted ester allowed expansion of ester diversity *via* late-stage functionalisation.

The presence of the C-7 Weinreb amide allowed further functionalisation by introducing either C-7 acyl substituents using Grignard reagents or C-7 amides using transamination. Successful C-7 acylation of Weinreb amides was achieved using aryl, alkyl and alkenyl Grignard reagents. High chemoselectivity at the side chain (Tables 1–3 and Scheme 4), with no reactions at the C-5 esters, was observed, except in the Grignard addition to 38iv which gave some C-5 ketone formation leading to 40 (Scheme 6). The high chemoselectivity of Grignard addition was attributed to both the hindered nature of the bicyclic tetramate system and the stability of a magnesium-chelated intermediate with the Weinreb amide, which also avoided over-addition leading to tertiary alcohols.<sup>59</sup> Since it has been reported that this Mg-chelated intermediate was only stable at low temperature, the process required a low-temperature reaction quench.<sup>59</sup> In the synthesised tetramate ketones, the *cis* relative stereochemistry at C-2/C-5 was supported by NOE analysis (Fig. S2, ESI†). With consistent H-2 chemical shifts across the series, *cis*-2,5 stereochemistry was assumed to be conserved from the starting tetramate Weinreb amides (Tables 1–3).

Tautomers existed for tetramates 41–73 as evidenced by the presence of two H-2 chemical shifts for each tetramate and the distinctive two sets of carbonyl chemical shifts across the series (Tables 1–3). The assignment of tautomeric forms was assisted

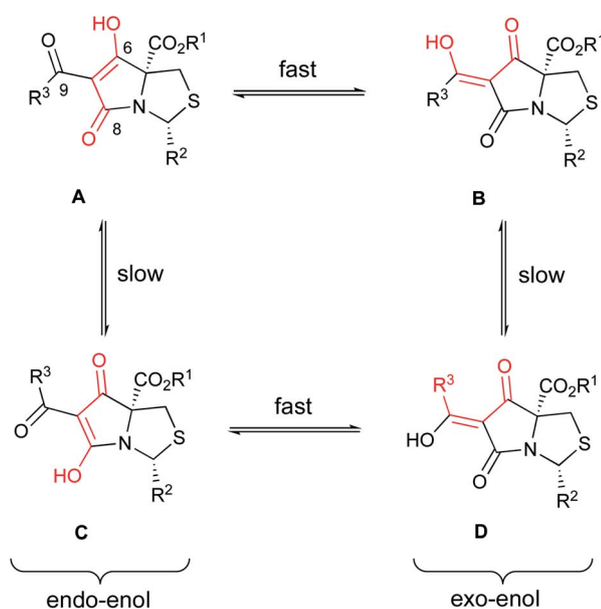
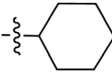
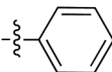
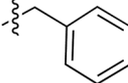
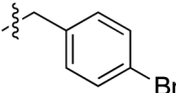
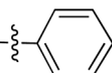
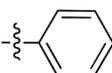
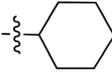
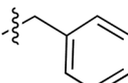


Fig. 3 Tautomeric behaviour of tetramate ketones.



Table 1 Yield, H-2, C-6, C-8 and C-9 <sup>1</sup>H and <sup>13</sup>C chemical shifts (CDCl<sub>3</sub>) for tetramate ketones 41–51

	Grignard R <sup>3</sup> MgBr	R <sup>1</sup> =	R <sup>2</sup> =	δ (ppm)				AB/CD	Yield (%)
				H-2	C-6	C-8	C-9		
41				6.42	192.3	175.6	187.3	4.1 : 1	71
				6.47	—	—	—		
42				6.42	192.3	175.8	187.3	3.7 : 1	45
				6.46	—	—	—		
43				6.41	192.0	175.9	187.6	4.2 : 1	40
				6.46	—	—	—		
44				6.40	191.9	175.8	187.7	4.5 : 1	80
				6.44	—	—	—		
45				6.40	192.3	175.9	187.3	4.2 : 1	43
				6.44	—	—	—		
46				6.40	192.5	175.9	187.3	3.3 : 1	42
				6.42	—	—	—		
47				6.49	185.9	177.7	182.9	CD	33
48				6.49	189.9	177.1	181.5	0.2 : 1	46
				6.40	185.9	177.9	182.9		
49				6.49	186.2	177.9	182.6	CD	62
50				6.48	186.4	177.9	182.5	CD	53
51				6.47	185.9	178.0	182.9	CD	30

by HMBC analysis (Fig. S2, ESI<sup>†</sup>) allowing assignment of C-6, C-8 and C-9 carbonyl shifts, which indicated that the *Z-exo-enolic* D as the major or exclusive tautomeric form in the R<sup>3</sup> = phenyl, phenethyl and styrenyl series and that the *E-exo-enolic* form B as the major tautomer in the R<sup>3</sup> = ethyl and cyclohexyl series (Fig. 3), was favoured. For the major forms, tetramate ketones with R<sup>3</sup> = ethyl and cyclohexyl consistently exhibited more downfield C-6 chemical shifts than the R<sup>3</sup> = phenyl, phenethyl and styrenyl series (Tables 1–3), which supported the assignment of the former as tautomeric form B and the latter as form D. The C-6 chemical shifts in tetramate Weinreb amides 31–39 assigned as *endo-enolic* form A (Table S4, ESI<sup>†</sup>) were the lowest of C-6 chemical shifts among tetramate ketones. C-8 and C-9 chemical shifts of tautomeric pair BD were very similar, with C-8 being generally more downfield in D because of hydrogen bonding. C-9 Chemical shifts in the R<sup>3</sup> = styrenyl series were

more upfield than others due to the extended conjugation system of the styrenyl side chain. The trend observed in carbonyl chemical shifts and the assignment of tautomeric forms was consistent to those in previously synthesised tetramate ketones with a C-2 *t*-butyl group. Besides the nature of the solvents used for NMR analysis,<sup>29,31</sup> the ratio of tautomeric forms was also dependent on the nature of the C-9 ketone substituent.

To expand the scope of this process, heterocyclic Grignard reagents such as pyridyl and thiazolyl systems were investigated for Grignard addition. Pyridyl magnesium bromide was prepared with ethyl bromide as an entrainment agent and successful Grignard addition was achieved to give 74 along with 42 (Scheme 7). Although the yield of the Grignard reaction is reported to be increased by raising the ratio of ethyl bromide to 3-bromopyridine to 3 : 1,<sup>60</sup> this proved not to be optimal with



Table 2 Yield, H-2, C-6, C-8, C-9 <sup>1</sup>H and <sup>13</sup>C chemical shifts (CDCl<sub>3</sub>) for tetramate ketones 52–65

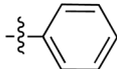
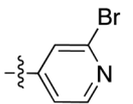
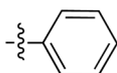
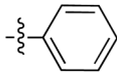
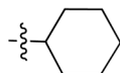
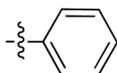
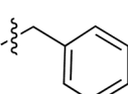
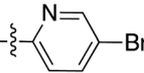
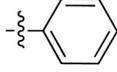
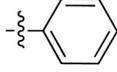
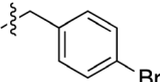
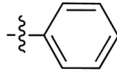
	Grignard R <sup>3</sup> MgBr	R <sup>1</sup> =	R <sup>2</sup> =	δ (ppm)				AB/CD	%Yield
				H-2	C-6	C-8	C-9		
52				6.41 6.44	195.3 —	176.1 —	187.1 182.4	3.8 : 1	35
53				6.28 6.30	195.9 —	176.4 —	186.3 —	3.3 : 1	33
54				6.41 6.44	195.3 —	176.3 —	187.1 —	4.0 : 1	39
55				6.40 6.43	195.1 —	176.4 —	187.4 —	4.0 : 1	30
56				6.39 6.42	195.0 —	176.4 —	187.6 —	4.3 : 1	57
57				6.35 6.38	195.7 194.3	176.8 172.9	187.0 186.0	3.7 : 1	29
58				6.38 6.42	195.3 —	176.4 —	187.1 —	4.6 : 1	37
59				6.38 6.41	195.5 —	176.4 —	186.9 —	5.1 : 1	36
60				6.40	189.8	175.3	187.1	CD	40
61				6.41 6.40	191.5 189.8	174.7 175.5	188.1 187.2	0.8 : 1	33
62				6.41 6.40	— 189.5	— 175.6	— 187.5	0.1 : 1	45
63	Phenethyl			6.42 6.40	— 189.4	— 175.6	— 187.6	0.2 : 1	73
64				6.38	189.8	175.6	187.1	CD	40
65				6.43 6.38	— 190.1	— 175.6	— 187.1	0.2 : 1	31

tetramate Weinreb amides since full reaction selectivity was lost, and the C-9 ethyl tetramate ketone **42** was obtained as by-product in every reaction.

Another widely used method of preparing organo-magnesium reagents is through halogen–metal exchange,<sup>61</sup> and this was applied to the 2-bromothiazole at room temperature



Table 3 Yield, H-2, C-6, C-8, C-9  $^1\text{H}$  and  $^{13}\text{C}$  chemical shifts ( $\text{CDCl}_3$ ) for tetramate ketones 66–73

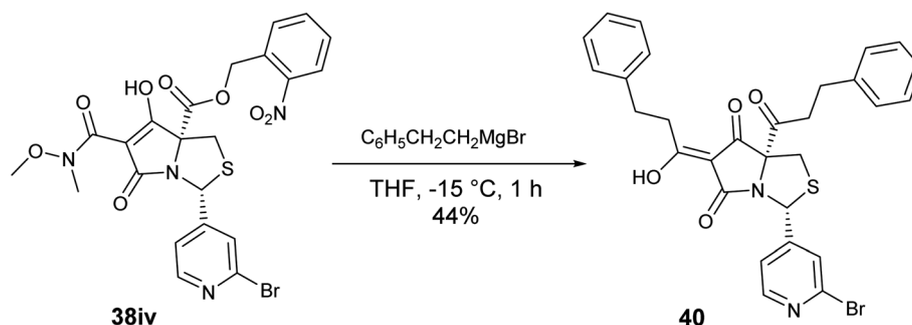
	Grignard $\text{R}^3\text{MgBr}$	$\text{R}^1 =$	$\text{R}^2 =$	$\delta$ (ppm)				AB/CD	%Yield
				H-2	C-6	C-8	C-9		
66				6.51	—	—	—	0.26 : 1	58
				6.46	187.5	176.0	176.2		
67				6.36	—	—	—	0.34 : 1	31
				6.32	186.7	176.3	176.6		
68				6.51	196.3	175.0	176.7	0.17 : 1	58
				6.46	187.5	176.1	176.2		
69	Styrenyl <sup>a</sup>			6.50	196.7	—	176.5	0.28 : 1	77
				6.46	187.7	176.0	176.3		
70				6.49	196.9	—	176.5	0.28 : 1	53
				6.45	187.9	175.9	176.3		
71				6.44	196.1	175.5	177.0	0.43 : 1	29
				6.40	187.4	176.5	176.7		
72				6.49	—	—	—	0.26 : 1	55
				6.44	187.4	176.2	176.3		
73				6.49	—	—	—	0.23 : 1	59
				6.44	187.3	176.2	176.3		

<sup>a</sup> The *E* stereochemistry of the styrenyl ketone pendant was confirmed by the *J* coupling value of H-1'' and H-2'' (consistently at *J* = 16.0 Hz, indicating the *trans* relationship between the two protons) on the side chain.

with  $i\text{Pr}_2\text{Mg}$ .<sup>62–64</sup> Subsequent Grignard addition to Weinreb amide **32iii** gave ketone formation but accompanied by hydrolysis and decarboxylation, giving derivative **75** (Scheme 8). *Exo*-enolic tautomeric form **D** was found for both tetramate ketones **74** and **75** (Table 4) based on comparison of the carbonyl shifts with the previously synthesised ketone series (Tables 1–3).

Along with the C-7 ketones **41–75** described above, C-7 carboxamides **76–94** could also be accessed from tetramate Weinreb amides **31–39** by transamination using the corresponding amines at reflux (Scheme 4). Interestingly, these

carboxamides exhibited two distinctive sets of H-2 shifts and carbonyl peaks for each compound, which were of approximately equal ratio (Table 5). The two sets of H-2 peaks were present in  $^1\text{H}$  NMR spectra of crude material, but did not arise after column chromatography or post-column acid wash. For **76**, 2D NOESY ruled out rotameric behaviour as the two sets of peaks did not show rapid dynamic exchange while 1D NOE analysis supported the same *cis*-relative stereochemistry at C-2 and C-5 (Fig. S2, ESI<sup>†</sup>). The assignment of carbonyl peaks was assisted by HMBC correlations, which showed that each of the



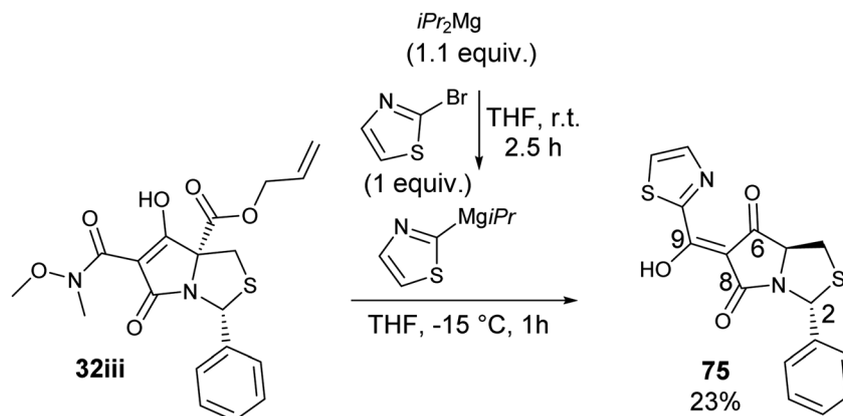
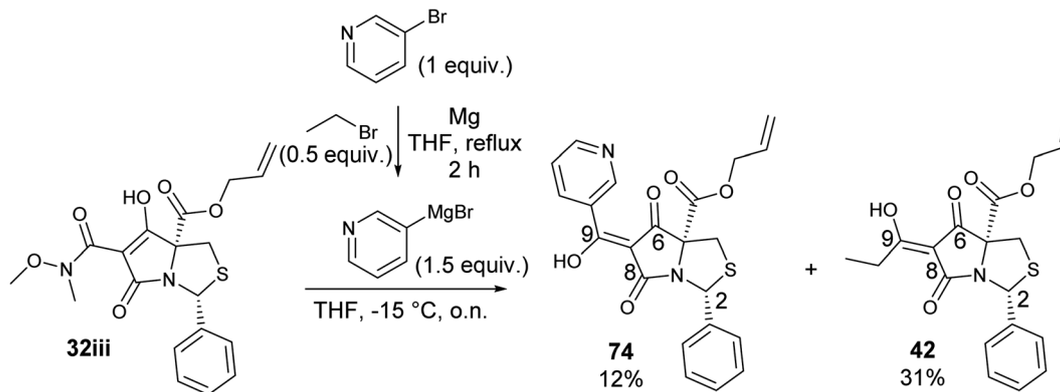


Table 4 Yield and H-2, C-6, C-8, C-9  $^1\text{H}$  and  $^{13}\text{C}$  chemical shifts ( $\text{CDCl}_3$ ) for 74 and 75

	Grignard $\text{R}^3\text{MgBr}$ or $\text{R}^3\text{Mg}i\text{Pr}$	$\text{R}^1 =$	$\text{R}^2 =$	$\delta$ (ppm)				AB/CD	Yield (%)
				H-2	C-6	C-8	C-9		
74				6.47	189.6	176.2	181.6	CD	12
75		—	—	6.48	189.2	173.0	177.9	CD	23

two H-2 shifts correlated with a different C-8 carbonyl (Fig. S2, ESI $^\dagger$ ). The only plausible explanation for the two sets of chemical shifts from a single reaction was tautomeric behaviour. The range of carbonyl chemical shifts was comparable and the consistent downfield trend in carbonyl peaks in tautomeric pair CD over AB was also similar to that observed with carboxamides. This tautomeric behaviour is also solvent dependent, so that, for example, for 88, in methanol, only one set of chemical shifts was observed in both  $^1\text{H}$  and  $^{13}\text{C}$  NMR spectra (Fig. S3C and D, ESI $^\dagger$ ), which appeared to be that of an averaged hybrid structure due to equilibration of the tautomeric form;

different behaviour was observed in  $\text{CDCl}_3$  (Fig. S3A and B, ESI $^\dagger$ ). Similar equilibration was observed in other polar solvents such as  $\text{CD}_3\text{CN}$  and  $\text{THF-}d_8$  but with broader signals, suggesting different rates of the solvent-assisted proton-transfer reactions.

#### Metal chelating properties of the synthesised tetramates

C-7 acylated tetramate ketones 41–75 were unsurprisingly metal-chelating, with their chelating abilities influenced by the bulkiness of both C-5 esters and C-9 ketone pendant groups. With the abundance of  $\text{Mg}^{2+}$  ions after the Grignard reactions,



Table 5 Yield, H-2, C-6, C-8, C-9 <sup>1</sup>H and <sup>13</sup>C chemical shifts (CDCl<sub>3</sub>) for tetramate carboxamides 76–94

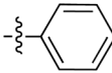
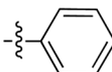
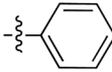
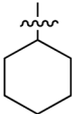
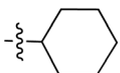
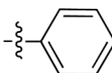
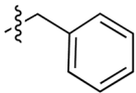
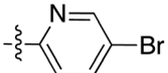
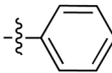
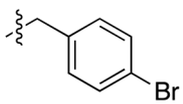
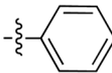
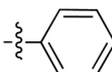
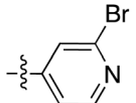
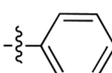
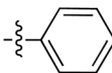
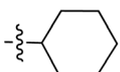
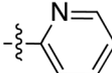
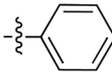
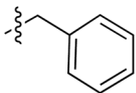
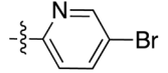
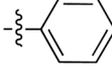
	Amine R <sup>3</sup> NH <sub>2</sub>	R <sup>1</sup> =	R <sup>2</sup> =	δ (ppm)				AB/CD	Yield (%)
				H-2	C-6	C-8	C-9		
76				6.27	187.1	173.2	165.8	1 : 1	30
				6.31	189.0	178.4	165.9		
77				6.26	187.0	173.3	165.7	0.9 : 1	48
				6.31	188.8	178.5	165.9		
78				6.26	187.4	173.4	165.7	1.1 : 1	49
				6.31	188.6	178.7	165.9		
79				6.25	187.5	173.4	165.7	1.1 : 1	61
				6.30	188.6	178.7	165.9		
80				6.24	187.1	173.5	165.8	0.9 : 1	58
				6.27	189.4	178.6	165.8		
81				6.25	187.1	173.5	165.8	0.9 : 1	35
				6.29	188.9	178.7	165.9		
82				6.25	186.9	173.4	165.8	0.9 : 1	47
				6.29	189.0	178.6	165.9		
83				6.26	187.0	173.4	166.7	0.9 : 1	56
				6.31	189.4	178.6	166.8		
84				6.14	186.6	173.5	166.7	1 : 1	52
				6.18	189.7	178.5	166.8		
85				6.26	187.0	172.6	166.7	1 : 1	46
				6.31	189.3	178.8	166.8		
86				6.24	187.2	173.5	166.5	1 : 1	44
				6.30	189.0	178.9	166.6		
87				6.31	187.4	173.6	166.4	1 : 1	33
				6.35	189.5	179.0	166.7		
88				6.25	187.3	173.5	166.4	1 : 1	57
				6.30	189.1	178.9	166.5		
89				6.23	187.0	173.6	166.7	0.9 : 1	34
				6.26	189.8	178.9	166.7		
90				6.23	187.0	173.6	166.6	1 : 1	67
				6.28	189.3	178.9	166.7		



Table 5 (Contd.)

	Amine R <sup>3</sup> NH <sub>2</sub>	R <sup>1</sup> =	R <sup>2</sup> =	$\delta$ (ppm)				AB/CD	Yield (%)
				H-2	C-6	C-8	C-9		
91			6.34	187.1	173.8	166.7	0.8 : 1	52	
			6.36	189.1	179.0				
92			6.14	186.5	173.5	166.7	1 : 1	43	
			6.18	189.7	178.5				
93			6.23	187.0	173.6	166.7	0.9 : 1	44	
			6.27	189.9	178.9				
94			6.24	186.9	173.6	166.7	0.9 : 1	47	
			6.29	189.4	178.8				

no meaningful product peaks could be detected in the NMR spectra of crude reaction mixture of **60**, for example, prior to acidic work-up (Fig. S4A, ESI<sup>†</sup>). However, once the crude material was washed with 0.5 M HCl, immediate sharpening of the resulting NMR spectrum was observed (Fig. S4B, ESI<sup>†</sup>). When subjected to flash column chromatography, the tetramate ketone again bound metal impurities from the silica, giving broad product peaks in the <sup>1</sup>H NMR spectrum (Fig. S4C, ESI<sup>†</sup>). Tetramate ketones were therefore routinely washed with 0.5 M HCl post-column to render them metal-free, which immediately sharpened the signals (Fig. S4D, ESI<sup>†</sup>). Similar

metal chelating behaviours were observed across the C-9 ethyl, phenethyl and styrenyl tetramate ketone series **41–73**. With bulkier C-5 esters, such as **65**, there was an improvement in the resolution of peaks post-column as compared to tetramates with smaller C-5 methyl esters (Fig. S4E and F, ESI<sup>†</sup>), supporting the hypothesis that steric hindrance posed by neighbouring C-5 esters reduced metal chelation. C-9 cyclohexyl ketones **52–56** demonstrated a similar broadness of peaks post-column chromatography but before acid wash. Interestingly, for tetramates **58** and **59** with bulkier C-5 benzyl or substituted benzyl esters, sharper peaks were detected on <sup>1</sup>H NMR spectroscopic

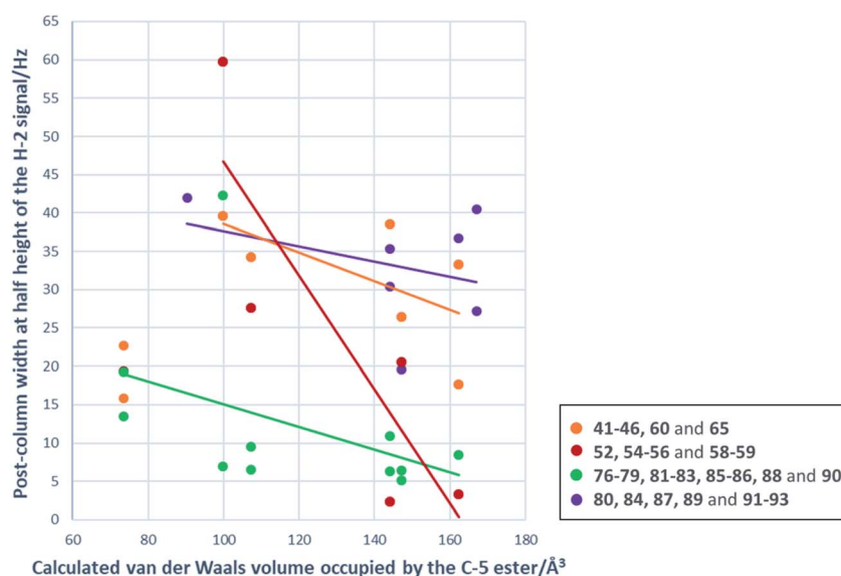


Fig. 4 Plot of post-column widths at half height/Hz for H-2 signals against van der Waals volume occupied by the respective C-5 ester. Subseries of compounds: orange – R<sup>3</sup> = ethyl or phenethyl, R<sup>2</sup> = phenyl; red – R<sup>3</sup> = cyclohexyl, R<sup>2</sup> = phenyl; green – R<sup>3</sup> = *N*-cyclohexyl or *N*-(1-adamantyl), R<sup>2</sup> = phenyl; purple – R<sup>3</sup> = *N*-cyclohexyl or *N*-(1-adamantyl), R<sup>2</sup> = pyridin-2-yl or bromo-pyridinyl.



examination of both the crude and post-column samples prior to acid wash (Fig. S5A, C and E, ESI<sup>†</sup>), consistent with reduced metal binding abilities in these compounds, although similar metal chelation behaviour was nonetheless discernible (Fig. S5C–F, ESI<sup>†</sup>). Comparison with tetramate ketones with the same C-5 ester also indicated the role played by the C-9 ketone pendant group in reducing metal binding. Thus, the presence of C-2 pyridyl rings in **57** and **71** broadened the post-column product peaks before acid wash, suggesting that the C-2 pyridyl rings could also participate in extended metal chelation (Fig. S6A, ESI<sup>†</sup>). As previously reported, tetramate ketones with a C-5 methyl ester required 2 M HCl solution to wash out metal ions,<sup>29</sup> and the fact that **57** and **71** could be washed free from metal ions with a much weaker 5% citric acid solution to give well-resolved and sharp <sup>1</sup>H NMR peaks suggested much reduced metal chelation resulting from the presence of a bulky C-5 ester (Fig. S6B, ESI<sup>†</sup>).

In order to investigate in more detail the types of metal ions potentially involved in chelation, metal-free tetramate ketone **60** was deliberately washed with 1 M aq. Ca(NO<sub>3</sub>)<sub>2</sub> solution (Fig. S7A, ESI<sup>†</sup>) and aq. 1 M FeSO<sub>4</sub> solution (Fig. S7B, ESI<sup>†</sup>). The extent of peak broadening suggested a higher affinity of metal chelation for Ca<sup>2+</sup> than Fe<sup>2+</sup>. Peak broadening induced by washing metal-free samples with aq. Ca(NO<sub>3</sub>)<sub>2</sub> solution has also been observed previously in tetramate ketones with C-2 *t*-butyl substituents. This observation in tetramate ketones was in contrast to tetramate esters and carboxamides, which demonstrated a preference for Fe<sup>2+</sup> with significant peak broadening (likely also due to paramagnetic behaviour) when washed with the same strength of metal solution.<sup>29</sup> This difference again

highlighted the structural dependency of metal chelation in tetramate derivatives. In addition, analysis by SFC-MS of a post-column sample of **63** prior to acid wash indicated the presence of *m/z* peaks that could be attributed to [M + Ca]<sup>2+</sup> and [M + Mg]<sup>2+</sup> but not [M + Fe]<sup>2+</sup> or [M + Fe]<sup>3+</sup> (Fig. S8, ESI<sup>†</sup>). The presence of Ca<sup>2+</sup> ions in post-column metal-chelated tetramate ketones was also previously indicated by XPS analysis.<sup>29</sup>

Tetramate carboxamides **76–94** were also similarly found to be metal-chelating, displaying broadness of product peaks post-column prior to acid wash (Fig. S9A, ESI<sup>†</sup>). Tetramate carboxamides were also routinely washed with 0.5 M HCl solution to remove metal ions post-column (Fig. S9B, ESI<sup>†</sup>), although 5% citric acid solution was used instead if pyridyl groups were present. For tetramate carboxamide **91** with a bulky C-5 nitrobenzyl ester, a 5% citric acid wash resulted in a better-resolved and sharp <sup>1</sup>H NMR spectrum, while the H-2 and H-5 signals of **95** still remained relatively broad after acidic wash as has been reported previously (Fig. S9C and D, ESI<sup>†</sup>).<sup>33</sup> The relative extent of metal binding with increasing bulkiness of C-5 ester was not discernible by visual comparison of <sup>1</sup>H NMR spectra. With an NH linker, the bulkiness of the C-9 amide pendant probably played a less important steric influence than the C-9 ketone pendant in reducing metal chelation.

A primary goal of this work was to modulate metal binding properties of the tetramates, and the concept behind chemical modifications of C-5 esters was to reduce metal chelation through steric influence of the neighbouring C-5 groups. In order to quantify the extent of metal chelation, peak broadening as measured by width-at-half-height (*W*<sub>1/2</sub>) values for the H-2 signals in post-column <sup>1</sup>H NMR spectra was used (Fig. 4 and

Table 6 Physicochemical properties of active tetramates.<sup>a</sup>

Cpd No.	MW/Da	clogP	clogD <sub>7.4</sub>	PSA/Å <sup>2</sup>	MSA/Å <sup>2</sup>	%PSA	HBD	HBA	MIC against		
									<i>Streptomyces</i>		MRSA
									μg ml <sup>-1</sup>	μg ml <sup>-1</sup>	μM
54	427.52	3.56	0.88	83.91	572.86	14.6	1	4	3.91	7.81	18.3
55	429.53	3.58	0.91	83.91	602.20	13.9	1	4	7.81	7.81	18.2
56	469.50	4.41	1.72	83.91	662.98	12.7	1	4	3.91	3.91	8.33
58	477.58	4.60	1.84	83.91	647.55	13.0	1	4	3.91	3.91	8.19
59	556.47	5.39	2.60	83.91	668.03	12.6	1	4	0.49	1.95	3.50
61	449.52	4.09	1.54	83.91	587.27	14.2	1	4	n.t.	0.98	2.18
62	451.54	4.11	1.58	83.91	616.53	13.6	1	4	n.t.	3.91	8.66
63	491.60	4.94	2.39	83.91	677.79	12.4	1	4	n.t.	0.98	1.99
64	499.59	5.13	2.51	83.91	662.21	12.7	1	4	n.t.	3.91	7.83
68	447.51	4.22	1.98	83.91	559.30	15.0	1	4	7.81	7.81	17.5
69	449.52	4.24	2.02	83.91	588.30	14.3	1	4	7.81	1.95	4.33
70	489.59	5.07	2.83	83.91	650.24	12.9	1	4	1.95	1.95	3.98
72	497.57	5.26	2.95	83.91	634.90	13.2	1	4	7.81	3.91	7.86
73	576.46	6.05	3.70	83.91	655.43	12.8	1	4	n.t.	0.98	1.70
79	484.61	2.79	-0.72	95.94	684.24	14.0	2	4	1.95	3.91	8.05
86	496.62	2.10	-1.42	95.94	680.97	14.1	2	4	1.95	7.81	15.7
87	537.68	1.89	-1.66	108.83	736.12	14.8	2	5	7.81	7.81	14.5
Ave.	484.26	4.20	1.51	86.79	640.40	13.6	1	4			

<sup>a</sup> MW was calculated by ChemDraw 19.1.1.32; clogP, clogD<sub>7.4</sub>, PSA, MSA, hydrogen bond donors (HBD) and acceptors (HBA) were calculated with Marvin (20.3.0), 2020, ChemAxon; %PSA was calculated by %PSA/MSA. (–) means tested but not active; n.t. means not tested. MIC values against MRSA that are <8 μg ml<sup>-1</sup> are highlighted in red.



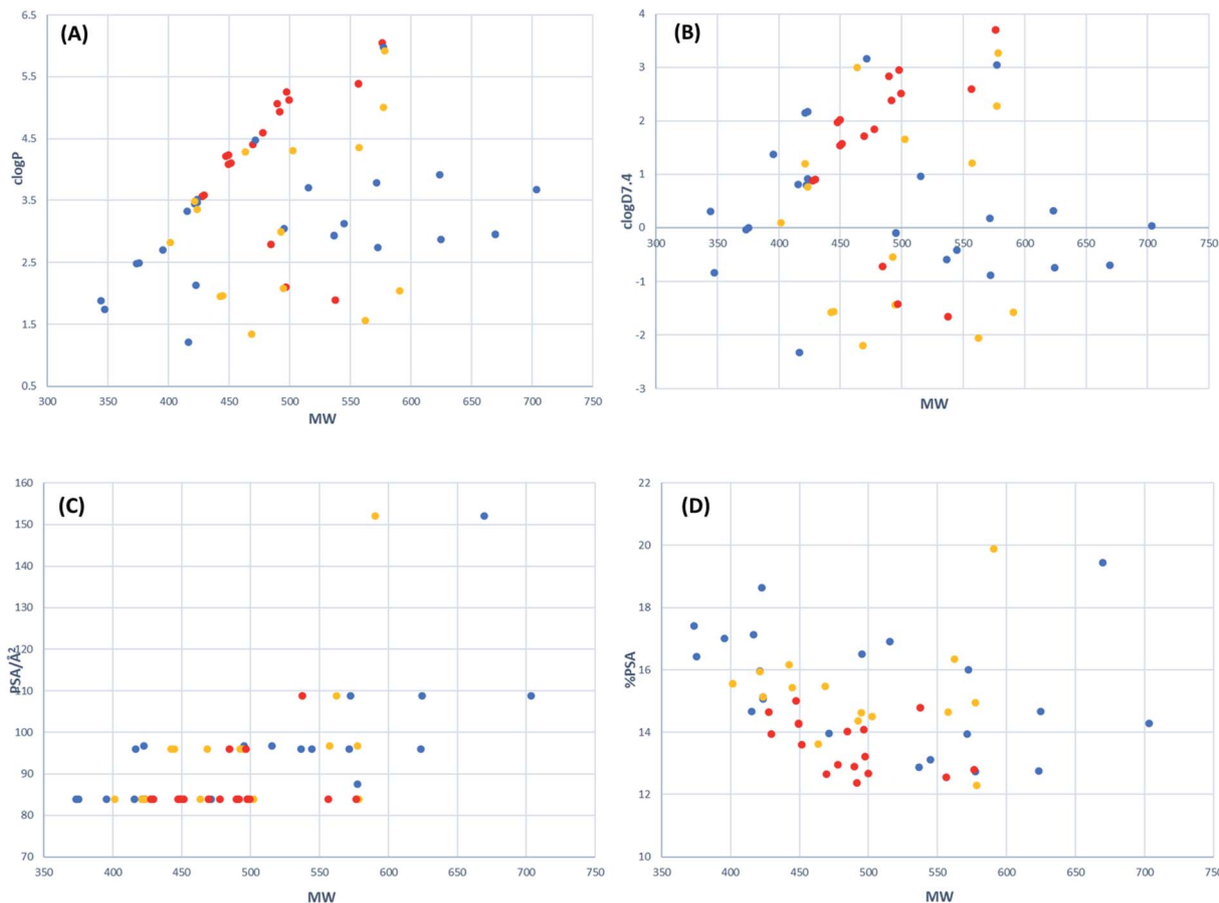


Fig. 5 Plots of (A)  $\text{clogP}$ , (B)  $\text{clogD}_{7.4}$ , (C)  $\text{PSA}/\text{\AA}^2$  and (D)  $\% \text{PSA}$  against MW/Da. Data representing the active tetramates are highlighted in red (MIC against MRSA,  $< 8 \mu\text{g ml}^{-1}$ ); tetramates exhibiting mild activity ( $8 \mu\text{g ml}^{-1} < \text{MIC against MRSA} \leq 32 \mu\text{g ml}^{-1}$ ) are represented in yellow; and tetramates with no activity are shown in blue.

Table S5,  $\text{ESI}^\dagger$ ). In cases where two sets of H-2 signals were present in the NMR spectrum of a single compound due to tautomeric behaviour, the major peak was taken for calculation. Moreover, since the principle behind chemical modifications of C-5 esters was to reduce metal chelation through steric influence of the neighbouring C-5 groups, their bulk was estimated by calculation of their van der Waals volumes ( $\text{\AA}^3$ ) and correlated with the post-column H-2 widths at half height (Fig. 4). Besides groups at the C-5 position, the metal chelation behaviour was also complicated by the identity of C-2 and C-9 pendants. Therefore, compounds were grouped into their respective sub-series and the trend of increasing C-5 steric bulkiness analysed separately. Across all series, an increase in the bulkiness of C-5 ester generally correlated with a decrease in peak broadening, suggesting reduced metal-binding ability and hence validating the working principle behind chemical modifications at C-5 esters (Fig. 4). Tetramates with C-2 heterocycles were generally more metal-binding, as heterocycles could also participate in extended metal chelation. The smallest extent of peak broadening among polar heterocycle-containing tetramate carboxamides was observed with **87** with C-2 pyridin-2-yl ring and C-5 cyclohexyl ester (Table S5,  $\text{ESI}^\dagger$ ). The best combination for reducing metal chelation was **58** with C-9 cyclohexyl ketone

pendant and C-5 benzyl ester, with a post-column H-2 width at 2.4 Hz, which is almost as sharp as non-metal binding tetramates.<sup>34</sup>

### Biological evaluation

Tetramate ketones **41–75** and carboxamides **76–94** were tested for their antibacterial activities against Gram-positive MRSA and *Streptomyces* and Gram-negative *E. coli* and the most active systems are reported in Table 6. No activity against *E. coli* was observed. Tetramate ketones **41–51** with C-9 ethyl or phenyl ketone pendants were not active while C-9 cyclohexyl, phenethyl and styrenyl ketone pendants **52–73** were more active against Gram-positive MRSA and *Streptomyces* (Table S6,  $\text{ESI}^\dagger$ ). Tetramate carboxamides **76–94** with C-9 cyclohexyl or adamantyl amide pendants also demonstrated activities against Gram-positive bacteria (Table S6,  $\text{ESI}^\dagger$ ). A key initial observation was that modification at C-5 esters did not result in activity loss, with bulkier C-5 esters actually enhancing the antibacterial potency against MRSA.

The drug-likeness of active tetramates was further assessed by calculation of their physicochemical properties, for which high oral availability, aqueous solubility and low protein plasma



Table 7 Cytotoxicity screen of selected cysteine-derived tetramates

Cpd no.	MIC ( $\mu\text{g ml}^{-1}$ ) against MRSA	IC <sub>50</sub> <sup>a</sup> ( $\mu\text{g ml}^{-1}$ ) against				MRSA selectivity <sup>b</sup>
		HeLa	HEK-239	CaCo	MDCK	
50	31.3	31.3	31.3	31.3	62.5	1.0–2.0
60	15.6	31.3	31.3	31.3	62.5	2.0–4.0
61	0.98	7.81	15.6	7.81	7.81	8.0–16.0
62	3.91	15.6	31.3	15.6	31.3	4.0–8.0
63	0.98	7.81	15.6	7.81	15.6	8.0–16.0
64	3.91	31.3	31.3	31.3	31.3	8.0

<sup>a</sup> IC<sub>50</sub> of 62.5–125  $\mu\text{g ml}^{-1}$  is considered non-toxic; IC<sub>50</sub> of 15.6–31.3  $\mu\text{g ml}^{-1}$  – less toxic and IC<sub>50</sub> < 8  $\mu\text{g ml}^{-1}$  – toxic. <sup>b</sup> The selectivity index for MRSA was calculated as IC<sub>50</sub>/MIC.

binding are considered to be important.<sup>65</sup> The parameters MW (Da), clogP, clogD<sub>7.4</sub>, polar surface area (PSA/Å<sup>2</sup>) and relative PSA (as a % of molecular surface area, MSA/Å<sup>2</sup>) of the synthesised tetramates were calculated (Table 6). clogP and clogD<sub>7.4</sub> predictions used Marvin 20.3.0, 2020, ChemAxon where the pool of fragments with assigned logP values is based on the data set of Viswanadhan *et al.*<sup>66</sup> While an MIC of 16–32  $\mu\text{g ml}^{-1}$  against tested pathogenic organism is considered a suitable minimum at a preliminary evaluation stage,<sup>67</sup> effective antibacterial activity of cysteine-derived tetramates synthesised in this project was defined by an MIC against MRSA of < 8  $\mu\text{g ml}^{-1}$ .<sup>68</sup> The ranges of physicochemical properties for active tetramate derivatives were MSA of 559–737 Å<sup>2</sup>, MW of 427–577 Da, clogP of 1.8–6.1, clogD<sub>7.4</sub> of –1.7–3.7, PSA of 83–109 Å<sup>2</sup> and relative PSA of 12–15%. These physicochemical boundaries are comparable to antibacterial active tetramate carboxamides previously reported (MSA of 530–600 Å<sup>2</sup>, clogP of 1.2–2.5, clogD<sub>7.4</sub> of –1.1–0, PSA of 65–80 Å<sup>2</sup> and relative PSA of 11–14%)<sup>30,69</sup> and these properties also resemble those of clinically used antibacterial fluoroquinolones (MSA of 380–550 Å<sup>2</sup>, clogP of 0.6–3.7, clogD<sub>7.4</sub> of –2.7–0.5, PSA of 60–120 Å<sup>2</sup> and relative PSA of 13–24%).<sup>68</sup> The chemical space occupied by the active tetramates was further delineated by plots of clogP, clogD<sub>7.4</sub>, PSA and relative PSA against MW (Fig. 5). For active tetramates, their clogP and clogD<sub>7.4</sub> values broadly correlate with MW while relative PSA changes in the inverse direction to MW (Fig. 5A, B and D). Such trends in structure–activity relationships were previously noted for antibacterial active tetramate carboxamides.<sup>69</sup> Most of the active tetramates possess a PSA of 82–84 Å<sup>2</sup> (Fig. 5C). The narrow ranges of PSA and relative PSA indicate the small variation in polarity which is required to maintain antibacterial activity against MRSA. Given that the work reported herein is synthesis-led, without the underpinning detailed mechanistic understanding of biological/biochemical mechanism, that the most active compounds occupy a small region of chemical space as defined by MW, clogP, PSA and % PSA (Fig. 5A–D) clearly highlights the benefit and role of chemical library synthesis and detailed evaluation in antibacterial drug discovery.

While some tetramates demonstrated good antibacterial activity against Gram-positive strains of bacteria including MRSA, no activity in Gram-negative *E. coli* was observed. This

observation was nevertheless in accordance with previous observations that Gram-positive and Gram-negative activities have different requirements of physicochemical properties.<sup>68</sup> It was previously reported that larger and more lipophilic C-7 acyltetramates with MW > 620 Da, clogP > 3, clogD<sub>7.4</sub> > 1, relative PSA < 12% were more easily transported by efflux pumps in Gram-negative *H. influenzae*.<sup>69</sup> Among tetramates that were tested active against MRSA, while none of their MW exceeds 620 Da and most of their relative PSA were > 12%, the majority of them have clogP > 3 and clogD<sub>7.4</sub> > 1, which increases their likelihood of being transported out by efflux pumps in Gram-negative bacteria. The trend of differential requirements between Gram-positive and Gram-negative activities was also noted by O'Shea and Moser where they reported the average physicochemical properties for selective Gram-negative activity as MW = 414, clogP = –0.1, clogD<sub>7.4</sub> = –2.8 and relative PSA = 42% and for selective Gram-positive activity as MW = 813, clogP = 2.1, clogD<sub>7.4</sub> = –0.2 and relative PSA = 30%.<sup>68</sup> Most

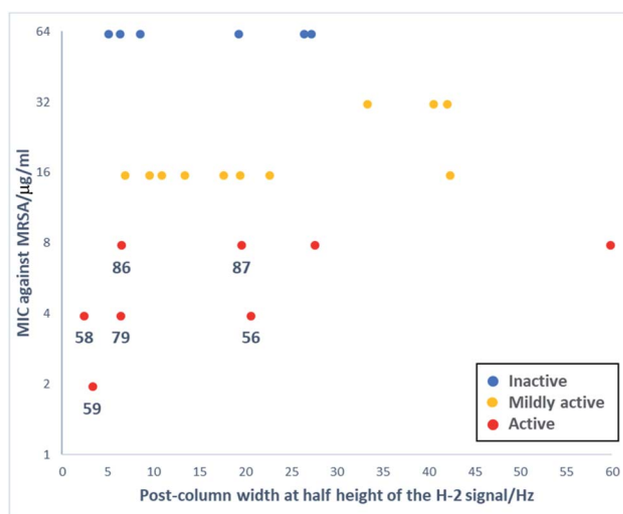


Fig. 6 Plot of MIC values against MRSA ( $\mu\text{g ml}^{-1}$ ) against post-column widths at half height of the H-2 signal (Hz) for C-5 modified tetramates. Data representing the active tetramate are highlighted in red (MIC against MRSA, < 8  $\mu\text{g ml}^{-1}$ ); tetramates exhibiting mild activity ( $8 \mu\text{g ml}^{-1}$  < MIC against MRSA  $\leq 32 \mu\text{g ml}^{-1}$ ) are represented in yellow; and tetramates with no activity are shown in blue.



tetramate-containing natural products exhibit selective activity against Gram-positive bacteria while only a few (such as epicoecarine A, zopfiellamide A and kibdelomycin) have been reported to be active against Gram-negative bacteria and mycobacteria; this has been attributed to their more complex cell wall and membrane structures which impede membrane permeability.<sup>70–77</sup> This suggests that tetramate-based derivatives generally occupy a chemical space more intrinsically in accordance with the requirement for Gram-positive activities. The average values of physicochemical properties for the active tetramates are MW = 484, clogP = 4.2, clogD<sub>7.4</sub> = 1.5 and relative PSA = 14% (Table 7), which fit better into the chemical space for selective Gram-positive activities as described by O'Shea and Moser.<sup>68</sup>

Interestingly, unlike C-6/C-9 modified tetramates,<sup>34</sup> reduced metal chelation ability in the C-5 modified tetramates did not result in a loss of antibacterial activity. Those C-5 modified tetramates that showed some antibacterial activity with MIC values against MRSA  $\leq 62.5 \mu\text{g ml}^{-1}$  were selected, and their MIC values plotted against post-column peak broadening ( $W_{1/2}$ /Hz of the H-2 signal) (Fig. 6). The absence of correlation between activity and peak broadening indicated that among tetramates with C-5 esters, antibacterial activity does not depend on metal chelation, with **56**, **58–59**, **79** and **86–87** showing reduced metal binding along with good antibacterial activity with MIC values against MRSA  $\leq 7.81 \mu\text{g ml}^{-1}$ .

Tetramate ketones **54**, **56**, **59** were further tested against MRSA in the presence of metal ions (culture medium containing Fe<sup>3+</sup>, Ca<sup>2+</sup>, Mg<sup>2+</sup> and Zn<sup>2+</sup> in the same concentrations as in blood) and in this case, antibacterial activity was retained, with MIC values of 7.81, 7.81 and 0.98 ( $\mu\text{g ml}^{-1}$ ) respectively. While there is evidence of reduced metal chelation in these tetramate systems, it clearly does not compromise their antibacterial activity, which is in contrast with our previous attempt to directly block metal complexation.<sup>34</sup> However, complete loss of activity in the presence of serum/blood was found, likely due to plasma protein binding due to relatively high lipophilicity (Table 7). This outcome highlights the challenge in antibacterial drug discovery of simultaneous multiparametric optimization, including consideration of at least efficacy, toxicity, membrane permeability and plasma binding.

The selectivity of tetramates **50** and **60–64** to prokaryotic bacteria cells over eukaryotic mammalian cells was analysed using cytotoxicity screens against HeLa, HEK-239, CaCo and MDCK cell lines. The selectivity index for MRSA was calculated as the ratio of IC<sub>50</sub> in respective cell lines against the MIC value (IC<sub>50</sub>/MIC) (Table 7). Tetramates **61–64** demonstrated enhanced selectivity towards MRSA, with **61** and **63** being more active with a MIC value of 0.98  $\mu\text{g ml}^{-1}$  and selectivity of more than 8-fold. While **62** and **64** were less active, they were also generally less toxic, with selectivity for MRSA as good as 8-fold. The different cytotoxicity towards the tested cell lines also suggested that there could be a viable therapeutic window for these tetramates even at high dose so long as specific tissue toxicities were also observed. It has been proposed that a reduction in toxicity could be achieved by increasing compound complexity with introduction of sp<sup>3</sup> hybridised carbon and reduction of aromatic

rings to improve specificity of target interaction.<sup>78,79</sup> In this regard, the chiral bicyclic tetramate template is an attractive scaffold and among the tetramates tested for cytotoxicity, and tetramate ketones **61** and **63** are particularly promising leads for further optimisation.

## Conclusion

The synthesis of cysteine-derived tetramate ketones and carboxamides with various C-5 esters, designed to decrease metal chelation by increasing bulkiness of the groups near the centre of metal binding, is reported. To achieve this goal, a library of L-cysteine esters was synthesised using acid-catalysed esterification of L-cystine in cyclohexane followed by selective reduction of the disulphide bond, with high yield and enantiopurity. This was followed by condensation with aldehydes and N-acylation, and a key step introduced in the synthetic route to incorporate a Weinreb amide to the N-acyl side chain of malonamides under Sucunza's acylation conditions allowed subsequent chemo-selective Dieckmann cyclisation with DBU to proceed in the direction that retained the C-5 ester and C-7 Weinreb amide. The resulting key intermediates, tetramate Weinreb amides, allowed further functionalisation to easily access C-7 ketones and carboxamides. The target tetramates were synthesised with a *cis* relationship between C-2 substituents and C-5 esters, as mixtures of tautomers in non-polar solvent such as CDCl<sub>3</sub>. Qualitative comparison of post-column <sup>1</sup>H NMR spectra suggested reduced metal binding in tetramates with bulkier C-5 esters and C-9 ketone pendants. Interestingly, modifications towards bulkier C-5 esters resulted in an enhancement of antibacterial activity against Gram-positive bacteria including MRSA, which indicated the potential for controlling metal chelation without compromising biological activity at C-5 position of the bicyclic tetramate system. Some synthesised tetramate ketones and carboxamides exhibited promising antibacterial potencies against Gram-positive MRSA with MIC values  $< 8 \mu\text{g ml}^{-1}$ , which were found to be independent from metal chelation. The antibacterial activity of these tetramates was maintained when tested in solvent media with metal ions of the same concentrations as in blood. Analysis of their physicochemical properties delineated a chemical space occupied by the active tetramates, with MSA of 559–737 Å<sup>2</sup>, MW of 427–577 Da, clogP of 1.8–6.1, clogD<sub>7.4</sub> of –1.7–3.7, PSA of 83–109 Å<sup>2</sup> and relative PSA of 12–15%. The structure–activity relationships indicated that clogP and clogD<sub>7.4</sub> of the active tetramates change approximately in proportion to MW while relative PSA changes in the inverse direction but there is only a narrow activity window for PSA variations. The active tetramate derivatives are generally Lipinski's rule compliant, which predict for good oral availability. The set of physicochemical properties fit to the chemical space of antibacterial drugs that possess selective Gram-positive activity. A subset of tetramates was tested for their cytotoxicity in mammalian cells and some tetramates exhibit good selectivity towards prokaryotic bacterial cells. Overall, the suitability of the natural product inspired bicyclic tetramate template as a promising structural motif for the development novel antibacterial drugs, with good anti-MRSA



potencies and appropriate physiochemical properties, to be drug-like, coupled with a potential for multi-targeting mechanisms and low eukaryotic cytotoxicity, has been demonstrated.

## Data availability

The datasets supporting this article have been uploaded as part of the ESI† material.

## Author contributions

RRZ conducted chemical synthesis, data collection and analysis, and manuscript preparation; MG, AP and DP conducted bioassays and reviewed the manuscript; MGM supervised the work and wrote the manuscript.

## Conflicts of interest

There are no conflicts to declare.

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