2-(HETEROATOM-SUBSTITUTED) METHYL PENEMS. I. SULPHUR DERIVATIVES

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stituted) alkylthio or heterocyclylthio radical are reviewed. The rationale at the basis of the programme is checked against the obtained microbiological data.

Following the discovery of a natural active structure, a great deal of research work has usually to be spent altering the original molecule in order to achieve optimal biological effects; this task is further aggravated when dealing with man-made structures, such as penerns. Even so, the poorness of consolidated structure-activity relationships inside the penem family doea appear surprising: while the importance of the hydroxyethyl chain at C-6 was **soon** recognized, the role of substituents at position 2 in modulating the antibacterial activity is still matter of conjecture. On the subject, it has been recently written that "..variation of the 2-substituent exerts only a modest effect on in vitro activity..", the main differences appearing rather "to reflect the effect [of the 1 substituent] an the lipophilicity of the molecule." A quite opposite rationale **was** adopted by us in approaching this field: **we** surmised that. **because** of the enamine system common to penems and cephalosporins, the inductive effect of the 2-position side chain would strongly influence the reactivity of the ß-lactam carbonyl and therefore be of the utmost relevance. This holds true for cephalosporins, where not only the average Gram-negative activity can be correlated to the inductive substituent constants for the C-3 side chain  $^2$ , but a heteroatom-substituted methyl in this position **3**  is usually more beneficial than a heteroatom attached directly to the nucleus : hence our particular interest in 2-CH<sub>2</sub>X (X= S,0,N) penem derivatives. We wish here to report our unpublished results in the **area.** and point out **some** discrepancies with the general'statement referred to above. The sulphur derivatives we considered can be divided in two families, I and II, according to whether the sulphur atom is further linked to a heterocyclyl or alkyl residue. The former family **was** pursued first, since it bears obvious relationships with the majority of cephalosporins present on the mar-*5*  ket, and preliminary studies on a representative lacking the C-6 side chain had been encouraging.



Initially, our synthetic plan closely duplicated Woodward's strategy;  $^6$  i.e. a phosphorane, already incorporating the heterocyclylthio moiety, **was** cyclized to each individual target compound. his route, which suffered from low yields owing to the sensitivenessof the thioester intermediates and to the problematic choice of the multiple protecting groups, had been expedient for obtaining a sample of the first member of the series, the methyltetrazolyl derivative Ia, but lacked the versatility and productivity required for the preparation of a consistent number of analogues. For this purpose **we** envisaged the possibility of exploiting a 2-hydroxymethylpenem as a late **common** interme-8 diate and, since an expressly designed carboryl protecting group had proved invaluable for the first synthesis of Ia, this common precursor was sought in the 1-phenoxyethyl ester 5a (Scheme I). To our end, glycolic acid **was** treated with p-nitrobenzyl chloroformate to give the carbonate (43%) and the dimeric derivative 1b; the former was converted into the thioacid 1c (EtOCOC1/NEt<sub>3</sub>,  $CH_2Cl_2$ ; then  $H_2S/NEt_3$ ) and used for acetate displacement on azetidinone 2a. The thioester 3a thus obtained was condensed with I-phenoxyethyl glyaxylate and then processed along the popular phopharane sequence<sup>6</sup> to give the fully protected 2-hydroxymethylpenem  $\frac{4a}{2a}$ ; the desired free alcohol  $\frac{5a}{2a}$  was thence unmasked in high yield by catalytic hydrogenation. The conversion of this compound into the thioether  $\frac{8a}{20}$ , a key intermediate in the first synthesis of Ia, clearly demonstrated the potentiality of hydroxymethylpenems  $5$  in the preparation of a large number of heterocyclylthiomethyl derivatives; such conversion was accomplished either by sequential mesylation  $\binom{CH_3SO_2CI/NEt_3}{2}$ ,  $\binom{CH_2CI_2}{2}$  and displacement with 5-mercapto-1-methyltetrazole (MMT sodium salt, THF, few hours, 0°C), or in a single step via the Mitsunobu reaction  $^{10}$  (MMT/PPh<sub>3</sub>/diethyl azodicarboxylate).

Having established the practicability of the new synthetic approach, the chemistry of the protecting ~FOUPG and of the exchange reaction **was** more attentively examined. First, since hydrolysis of the ester group on the final compounds **was** one major problem, **we** devised to perform this atep prior to thiol introduction. Thus, p-nitrobenzyl glyoxylate was substituted far the 1-phenoxyethyl analogue in the above sequence leading to  $\frac{4a}{\gamma}$ , thereby obtaining  $\frac{4b}{\gamma}$ ; catalytic hydrogenation in a biphase EtOAc-aq.NaHCO<sub>3</sub> system (the carbonate being removed first) then afforded the sodium salt 6 in satisfactory yield. A convenient temporary protection for the carboxyl function of 6 was found in the labile tert-butyldiphenylsilyl group, which **was** selectively introduced by performing the silylation in THF (in CH<sub>2</sub>C1<sub>2</sub> formation of the trisilylated compound 4c could not be avoided); the obtained di-<br>silylated intermediate 5b was then activated on the free hydroxyl (MsCl/NEt<sub>3</sub>, CH<sub>2</sub>C1<sub>2</sub>) and reacted<br>in situ with silylated intermediate 5b was then activated on the free hydroxyl (MsCl/NEt<sub>3</sub>, CH<sub>2</sub>Cl<sub>2</sub>) and reacted<br>in situ with the thiol to give the desired thioether  $\frac{\partial b}{\partial x}$ . Unfortunately, double desilylation of<br> $\frac{\partial b}{\partial x}$ in situ with the thiol to give the desired thioether 8b. Unfortunately, double desily lation of under the conditions of silyl removal at position 8 whenever the carboxyl is unmasked first. In Order to avoid this problem, selective activation of the primary alcohol **was** performed on the fully order to avoid this problem, selective activation of the primary alcohol was performed on the full<br>deprotected synthon 7. This material, easily accessible from azetidinone 2b via thioester 3b and<br>exact to accepting to a gr 11 deprotected synthon 7. This material, easily accessible from azetidinone 2b via thioester 3b and<br>penem 5c according to a previous methodology,<sup>11</sup> when temporarily protected as the TBDPS ester 5d<sup>12</sup><br>underwent meavlation a underwent mesylation and in situ displacement (MMT sodium salt/CaCO<sub>3</sub>, THF-CH<sub>2</sub>Cl<sub>2</sub>) to afford 9b and  $\sim$ <br>otected as the TBDPS ester 5d<sup>12</sup><br>3<sup>,</sup> THF-CH<sub>2</sub>C1<sub>2</sub>) to afford 9b and<br>closures of all ul esters appear thence  $I_{\alpha}$  (HOAc-THF-H<sub>2</sub>O, 3:1:1; 40 min,r.t.).<sup>13</sup> The extremely mild cleavage of allyl esters recently 14 introduced by McCombie et al. **came** welcome in this context; the allyl ester 5e often proved supe- **r"**  rior to 5a, 5c,<sup>15</sup> 5d as a precursor of 2-thiomethyl substituted penems, either through the mesylate

## SCHEME I



pNB= p-nitrobenzyl; TBDNS= SiMe  $_2^{\text{t}}$ , TBDPS= SiPh  $_2^{\text{t}}$ ; heterocycles A $\sim$ D are as shown in Table II

**OF** the Mitsunobu route.

Other obvious pathways to compounds I entailed the displacement with the heterocyclic thiol of either a halo or an acetoxy group, in analogy with methodologies routinary in the cephalosporin 16 either a halo or an acetoxy group, in analogy with methodologies routinary in the cephalosporin<br>field. However, reaction of MMT sodium salt with the acetate  $\frac{10a}{100}$  under a variety of conditions<br>gave, if any, only a acetates have been claimed to undergo easier displacement by nucleophiles than acetates,  $^{17}$  but reaction of MMT on 10b (from 5c and diketene) or on the sodium salt 10c (from 10b by hydrogenolysis)<br>was equally unrewarding. On the other hand, halomethylpenems proved to be reactive species, though plagued by stability problems. Compound 10d could be obtained by treatment of  $\frac{5}{3}$  with PPh  $\frac{3}{3}$  CCl  $\frac{4}{3}$ plagued by stability problems. Compound 10d could be obtained by treatment of 5c with PPh<sub>3</sub>/CCl<sub>4</sub><br>(24 h, r.t.), while 10e, accompanied by a consistent amount of 4-allyloxycarbonyl-5-chloromethyl-<br>
thiazele was secured fr (24 h, r.t.), while 10e, accompanied by a consistent amount of 4-allyloxycarbonyl-5-chloromethyl-<br>thiazole, was secured from reaction of 5e with SOC1<sub>2</sub> (pyr; CH<sub>2</sub>C1<sub>2</sub>-40°C) they both survived rapid<br>flash shumstasseship flash-chromatography, but yields were low and irreproducible. Better results were obtained from



halamethylpenems generated under milder conditions: in particular from the carbinols through a new 18 application of the Mitsunobu reaction, from **4-haloacetylthioazetidinones** by a low-temperature 19 20 Wittig condensation, from **3-bromomethyl-2-thiacephees** by ring-contraction with PPh at -40°C.0n 3 balance, however. the mesylate and Mitsunobu approaches remained the most satisfactory routes to penems  $I$ . Mesylates, e.g.  $10g$ , although unstable upon storage, could easily be reacted in situ and proved more handy than the corresponding tosylates, e.g. 10h. The Mitsunobu procedure turned out particularly appealing when **we** found that a modification in the order of addition, originally suggested by Volante<sup>21</sup> for the synthesis of thioesters, minimizes by-products and base-catalyzed side-22 reactions in this particular thioether synthesis. According to this methodology. diethyl azodicarboxylate and PPh<sub>3</sub> were first allowed to form a crystalline complex (THF, 30 min.r.t.), which then  $\overline{\phantom{a}}$ immediately reacted at  $0^{\circ}$ C when added to a THF solution of the carbinol  $\sum_{i=1}^{\infty}$  and the heterocyclic thiol. The functionalities present on **some** heterocyclic thiols had to be taken into account. The reaction The functionalities present on some heterocyclic thiols had to be taken into account. The reaction<br>of 8c to give the iminophosphorane 8d in the presence of excess Mitsunobu reagent is a singular<br>example <sup>23</sup> although in th 23 example, although in the instance the free amino group could be restored by mild acidic hydrolysis (THF-HOAc-H<sub>2</sub>O 3:1:1, overnight). Acid hydroxy groups had to be protected before condensation; in order to avoid any unnecessary lenghtening of the synthetic sequence, ally1 and silyl protecting groups **were** chosen for the purpose. Thus. the tetrazolylprogionic acid **was** protected as its

allyl ester 11b, (neat allyl alcohol, cat. 37% aq.HCl), while the thiadiazole derivative 12a was



protected as the <u>tert</u>-butyldiphenylsilyl ester 12b (TBDPS chloride, NEt<sub>3</sub>; THF). The difference in  $\mathbf{p} \in \mathbb{C}$  is the text-butyldiphenylsilyl ester 12b (TBDPS chloride, NEt<sub>3</sub>; THF). The difference in acidity between the two isomeric triazinones 13a and  $\frac{2^5}{4}$  was reflected by their need for protection, the l acidity between the two isomeric triazinones 13a and  $14^{25}$  was reflected by their need for protection: the less acidic isomer  $14$  readily reacted as such, while the strongly acidic hydroxyl group tion: the less acidic isomer 14 readily reacted as such, while the strongly acidic hydroxyl group<br>of 13a had to be protected as its silyl ether 13b. This last compound suffered desilylation upon attempted mesylate displacement, but could be condensed either with 5b or 5e under the previously reported Mitsunobu conditions.

A major problem in the preparation of thiomethylpenems I, II was their propensity to undergo equilibration to thiomethylenepenams III, IV. Shift of the double bond between the endo and the exo position following base treatment was first observed by Woodwa~d,~~ and it has since found **some**  28<br>utility for the synthesis of thia-analogs of clavulanic acid,  $^{27}$  for 2-oxopenam intermediates and also for obtaining 2-alkylpenems from alkylidenepenarn~~~ in cases where, owing to the **lack** of conjugation, the latter are the thermodynamically less stable isomers. For heterocyclylthiomethylpenem esters  $I$  the endo-exo equilibrium lies somewhere in the proximity of 30:70, and this proportion is even more unfavourable for the alkylthio analogs II, so that the success of the synthesis **was** largely dependent on the kinetic factor, **i.e.** on the ingenuity exerted in avoiding prolonged basic conditions during the exchange-deprotection sequence. The synthesis of **Ic** is illustrative. Condensation of 5<sub>g</sub> with 1-(2-dimethylaminoethyl)tetrazole-5-thiol (mesylate route) and quenching with HOAc afforded a 1:1 mixture of the endo/exo products, &e and 15a. In the presence of a large excess of acetic acid, this mixture underwent desilylation (TBAF, THF overnight) without signifi-<br>cant change in the isomer ratio, and then the desired intermediate 9e could be separated from the





## 15



slower-running exo-isomer 15b by chromatography on silanized silica. By using HOAc instead of sodium ethylhexanoate, deallylation of 9e with the Pd-PPh<sub>3</sub> complex furnished an inseparable 3:2 mixture<br>ethylhexanoate, deallylation of 9e with the Pd-PPh<sub>3</sub> complex furnished an inseparable 3:2 mixture<br>as *is and III* (and ins slower-running exo-isomer 15b by chromatography on silanized silica, by using HOAc instead of sodium<br>ethylhexanoate, deallylation of 9e with the Pd-PPh<sub>3</sub> complex furnished an inseparable 3:2 mixture<br>of Ic and IIIc (as the of Ic and IIIc (as the internal salts); under the usual conditions the sodium salt of the latter was exclusively obtained. In this instance, and whenever penams III were the predominant final products, it proved difficult to revert the **ex0** form into the equilibrium mixture in acceptable yields, presumably owing to the limited intrinsic stability of compounds I. Nevertheless, direct evidence of a reversible equilibrium being at work **was** often apparent in protected intermediates when the two forms happened to be easily differentiated by chromatography: for example on separate desilylatian of pure  $\frac{\partial f}{\partial x}$  and 15c the same mixture (ca. 1:1) of isomers 9f, 15d was obtained. Alkylidenepenams arising from base-catalyzed isomerization of penems are expected to possess the **more** stable configuration at the centres involved in the equilibration, i.e. to be *S* at C-3 and Z at the **ex0** double bond. Indeed, a single isomer had always been found in previous works 26.29 and assigned this structure. In one instance, however, we were able to detect by nmr spectroscopy the presence of a minor component (1:9), 15f, accompanying the major compound, 15e; they were considered, respectively, as the <u>E</u>, Z alkene isomers of the 8R, 6S, 5R, 3S form. Although the relative assignment **was** made by inference.30 the high deshielding experienced by the C-3 protons in both compounds, and the minimal mutual difference  $(\Delta \delta = 0.07$  ppm), ruled out an alternative interpretation in terms of a diastereomeric relationship at C-3.  $^{\rm 31}$ 

Another unusual alkylidenepenam structure was detected in the reaction mixture from 5e and 2mercapto-5-methyl-1,3,4-thiadiazole (mesylate route); it manifestly incorporated two elements of the heterocyclic thiol and on the basis of analytical and spectral data **was** formulated **as** 16a. ts of<br> $16a$ .<br> $(6a)^{33}$ This material, as well as any other alkylidenepenam that could be isolated reasonably free  $(\leqslant 5%)$ from its penem isomer, was conventionally deprotected, and the obtained salts (16b, IIIc-h; Table III) were assayed for antimicrobial activity (Table IV).

While the heterocyclylthiomethylpenems <sub>I</sub> are reminiscent of cephalosporins, the alkylthiomethyl analogs II offer different reasons for interest. In particular, compound IIa bears strict relationships with thienamycin and its penem counterpart. Sankyo's "1-thiathienamycin",  $\frac{34}{20}$  Compound IIb carries the carbamoyloxy functionality characterizing Farmitalia Carlo Erba's FCE 22101, $^\mathrm{11}$  and is the 2-homolog of another 1-thiathienamycin derivative,  $\stackrel{\text{Vb}}{\sim}$  the new Schering's candidate (Sch34343). $^{35}$ 



For the synthesis of compounds II the hydroxymethylpenem approach proved of very limited value, owing to the weak acidity of alkylthiols and to the sensitiveness of penems towards nucleophiles. Thus, the mesylate from 5g did not react in the presence of triethylamine with the protected

SCHEME I I



cysteamine  $17a$ , while reaction with the lithium salt of the latter (from the thiol and BuLi) resulted in  $B$ -lactam opening. Similarly, the carbinol 5e was decomposed by treatment with  $PBu_{3}^$ resulted in B-lactam opening. Similarly, the carbinol 5e was decomposed by treatment with  $PBu_3$ -<br> $Et_2S_2$  (with PPh<sub>3</sub> no reaction occurred), and condensation of 5e with 17a under Mitsunobu-Volante<br>conditions sous a mixt Et<sub>2</sub>S<sub>2</sub> (with PPh<sub>3</sub> no reaction occurred), and condensation of 5e with 17a under Mitsunobu-Volante<br>conditions gave a mixture of penem 22a and penam 23a in moderate yields. We therefore resorted to<br>the approaching of con the preparation of azetidinyl thioesters already incorporating the desired functionality. Following the preparation of azetidinyl thioesters already incorporating the desired functionality. Followi<br>an original Farmitalia route,  $36$  the 1,2-secopenicillanate 18a, obtained by trapping a penicillin-<br>deviced subthacia asid an original Farmitalia route,<sup>36</sup> the 1,2-secopenicillanate 18a, obtained by trapping a penicilli<br>derived sulphenic acid with propargyl alcohol, was converted into the bromo-sulphide 18b; this<br>----------------------------monocyclic &lactam, d~fferent from the activated hydraxymethylpenems, could **be** easily thialated derived sulphenic acid with propargyl alcohol, was converted into the bromo-sulphide 18b; this<br>monocyclic B-lactam, different from the activated hydroxymethylpenems, could be easily thiolated<br>with the lithium salt of 17a 20d was eventually obtained from the latter by sequential ozonolysis to 20b (80%), methanolysis on silica to 20c (30%) and sulphoxide reduction (PBr<sub>3</sub>. DMF-CH<sub>2</sub>C1<sub>2</sub>, 90%), any attempt to convert this 2 20c (30%) and sulphoxide reduction (PBr<sub>3</sub>. DMF-CH<sub>2</sub>C1<sub>2</sub>, 90%), any attempt to convert this 2 20 20 20 20 20 20 2 silica to 20c (30%) and sulphoxide reduction (PBr<sub>3</sub>, DMF-CH<sub>2</sub>Cl<sub>2</sub>, 90%), any attempt to convert this compound or its oxalimido precursor 20b into a triphenyl or triethoxyphosphoranylidene interme-<br>distance with unexpect compound or its oxalimido precursor 20b into a triphenyl or triethoxyphosphoranylidene interme-<br>diate met with unexpected failure. Phosphoranes 19 proved better suited than azetidinones 20 for<br>the ludinosity of any site of diate met with unexpected failure. Phosphoranes 19 proved better suited than azetidinones 20 for<br>the derivation of penem IIa. Thus 19b and 19c, easily obtained from the silylated parents, e.g.<br>10. <sup>37</sup> uses estimated with the derivation of penem IIa. Thus 19b and 19c, easily obtained from the silylated parents, e.g.<br>19a,<sup>37</sup> were activated either as the mesylates or the bromides and reacted with the protected cyste-<br>19a,<sup>37</sup> were activated man and the numerical contract of  $\frac{1}{2}$ ,  $\frac{1}{2}$  and  $\frac{1}{2}$ ,  $\frac{1}{2}$  and  $\frac{1}{2}$ ,  $\frac{1}{2}$  and  $\frac{1}{2}$  considers the interpreted cyst and  $\frac{1}{2}$  (lithium salt). Thermal cyclization of 2lb, followed by c resulting tris-protected penem 22b, finally afforded a sample of the desired 2-homo-1-thiathienaamine 17a (lithium salt). Thermal cyclization of 21b, followed by catalytic hydrogenation of the<br>resulting tris-protected penem 22b, finally afforded a sample of the desired 2-homo-1-thiathiena-<br>mycin IIa. Similarly, we co get in this series, IIb. The required thioacid 17d was synthesized from 2-mercaptoethanol by sequential monosilylation (TBDMS chloride/imidazole; DMF, overnight, quantitative), alkylation with chloroacetic acid (KOBu<sup>t</sup> 2 mol eq.; EtOH, 2h, 95%), activation<sup>38</sup> of the resulting acid 17c as the chloride (oxalyl chloride/NEt<sub>3</sub>; CH<sub>2</sub>Cl<sub>2</sub>.-15°C) and in situ treatment with H<sub>2</sub>S in the presence of a further amount of NEt<sub>3</sub> (30 min; 20%).<sup>39</sup> Reaction of said thioacid with the acetoxyazetidinone 2a, follower the conventio amount of NEt. (30 min; 20%).  $\tilde{ }$  Reaction of said thioacid with the acetoxyazetidinone 2a, followed amount of NEt<sub>3</sub> (30 min; 20%).<sup>39</sup> Reaction of said thioacid with the acetoxyazetidinone 2a, followed<br>by the conventional phosphorane build-up sequence, uneventfully yielded 21c and thence 22c after<br>brief beating in refl by the conventional phosphorane build-up sequence, uneventfully yielded 21c and thence 22c after<br>brief heating in refluxing toluene. Although penem 22d could be gained virtually uncontaminated from the exomethylene isomer 23d by selective unmasking of the primary alcohol under mild acidic conditions (THF/HOAc/H<sub>2</sub>O 2:1:1; 40 h, 80%), <sup>40</sup> the particular propensity of alkylthiomethylpenems to undergo double bond migration was apparent in the cleavage of the secondary silyl ether (Bu<sub>A</sub>NF.3H<sub>2</sub>O, HOAc-THF) on the adduct from 22d and trichloroacetyl isocyanate, whereupon the alkylidenepenam 23g and thence the sodium salt  $\frac{1}{2}$  were exclusively obtained.

The final adjustment of the synthetic sequence embodied a non-isomerative deblocking of protecting groups and the use of a more stable thioacid. Substitution of the TBDPS group for TBDMS in the route leading to 17d from 2-mercaptoethanol conveniently afforded 17e, which was then allowed to react groups and the use of a more stable thioacid. Substitution of the TBDPS group for TBDMS in the rolleading to 17d from 2-mercaptoethanol conveniently afforded 17e, which was then allowed to react with azetidinone 2b and co with azetidinone 2b and conventionally processed to the phosphorane 21f. Reversal of the cycliza-<br>tion-desilylation previously exploited <sup>41</sup> gave 21g and thence the penem carbinol 22g, uncontaminated<br>hy any penem isomen. by any penam isomer. Reaction of the latter with trichloroacetyl isocyanate (CH  $_2^{\rm C1}$   $_2^{\rm P}$ , -20 °C) followed by cleavage of the intermediate urethane (MeOH/SiO $_2$  230-400 Mesh; EtOAc, 5 h) afforded the carbamate 22h and finally, after deblocking of the pNB groups  $(Fe/NH_4Cl; THF-H_2O, 1 h)$ , penem IIb was secured.

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Table I - Spectral data of key intermediates

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## Table I - Continued



1) Film unless otherwise stated; 2) in CDC1<sub>3</sub> unless otherwise stated; 3) additional data: 1a:<br>mp 97-98° (isopropyl ether); 5g: [a]<sub>0</sub> +66°(1% CHC1<sub>3</sub>),  $\lambda_{max}$  (CH<sub>2</sub>C1<sub>2</sub>) 269 ( $\epsilon = 17,000$ ) and 323 nm<br>( $\epsilon = 6,800$ ); <u></u>



Table II - Spectral data of thiomethylpenems I, II

Table II - Continued

Im $\sim$	M	3420 3300, 3180. 1765, 1620, 1560	320	3.92(1H, dd), 4.42(2H,s), 5.64(1H,d)
In $\sim$	N	2250.1765.1615	259,316	$3.23(2H, t)$ , $3.84(1H, dd, 1.4$ and $6.0Hz$ , $4.54$ $(2H, ABq, 14.2Hz), 4.8(2H, t), 5.58(1H, d)$
$I \circ^*$ ∼	0	3420, 1760, 1700, 1610.1580	304( $s = 8.600$ )	
$\mathfrak{P}^*$	P	3600-3200, 1765. 1580, 1390	315	
$\stackrel{\text{Id}^*}{\sim}$	$\mathsf Q$	1765, 1605, 1570	$239($ $\varepsilon = 18.930)$ , $312($ $\varepsilon = 9,074)$	4.06(1H, dd, 1.5 and 6.1Hz), 4.70 and 5.00 $(2H, each d, 12Hz)$ , 5.64 $(1H, d)$ , 7.72 and 8.42 (2H, each d, 9.6Hz)
Џа*		$CH_2CH_2NH_2$ 3420, 1770, 1610	308	2.83 and 3.24 (each $2H, t, 6.7Hz$ ), 3.88 (1H, dd, 1.4 and $6.3Hz$ , $4.01(2H, ABq, 14.7Hz)$ , $5.64$ (1H,d)
		IIb* CH <sub>2</sub> CH <sub>2</sub> OCONH <sub>2</sub> 3600-3200,1765, 1610, 1570	310	

1) Potassium salt unless otherwise indicated: \*) Na<sup>+</sup> salt; \*\*) inner salt.



3) Data relative to  $H_5$ ,  $H_6$ , 2-CH<sub>2</sub> and R protons; HOD at 4.81 ppm as internal standard.

Compd.	R	ir (KBr) (cm ⊡ $\boldsymbol{v}$ max	$uv$ (H <sub>2</sub> 0) $\lambda_{\text{max}}(\text{nm})$	$H$ nmr $(D_{\gamma}0)$ $\delta$ (ppm)
IIIc* $\sim$	c	1760, 1610		$2.63(6H,s)$ , $3.35(2H,t)$ , $3.58(1H,dd,1.5$ and $7Hz$ , 5.00(2H,t), 5.40(1H,d,1Hz), 5.47(1H, $d, 1.5Hz$ , 6.44(1H.d.1Hz)
IIId $\sim$	D	1760,1610	$267($ $\varepsilon = 7,814)$ , 326( $\varepsilon = 4,787$ )	$3.55(1H, dd, 1.3 and 5Hz)$ , $5.43(2H,m)$ , 6.65 $(1H, d, 1Hz)$ , 8.3-8.6(3H.m)

Table III - Spectral data of thiomethylenepenams III, IV, 16b





1), 2) See footnotes 1, 2, Table II.

3) Data refer to  $H_3$ ,  $H_5$ ,  $H_6$ , vinyl and R protons; HOD at 4.81 ppm as internal standard.

Table IV - Antibacterial in vitro activities<sup>1,2</sup> of selected penems and penams



1) MIC values are given in µg/ml, and were determined in Isosensitest broth.

2) Organisms included in this Table are: S.a.S., Staphylococcus aureus Smith; S.p., Streptococcus pyogenes C203; E.c., Escherichia coli G; K.a. +, Klebsiella aerogenes 1082 E (producer of chromosomally mediated ß-lactamase).

3) Heterocycles A, C, D, H, O, Q are as indicated in Table II.

Table IY shows the minimal inhibitory concentration (MIC) values against four significative bacterial strains for the functionalized alkylthiomethyl penems IIa, IIb, and for a representative of 42 rial strains for the functionalized alkylthiomethyl penems  $\frac{IIA}{N}$ , IIb, and for a representative of<br>the heterocyclylthiomethyl family, <sup>42</sup> the methyltetrazole product Ia; the 2-(unsubstituted methyl) penem 10f was taken as a reference standard. These data are striking when compared to the early report by Bristol-Myers<br>report by Bristol-Myers' on 6-unsubstituted penems, which in median activity against Gram-negative bacteria favoured the 2-methylthiomethyl product over the 2-methyl reference, and the latter over the tetrazolylthiomethyl analogue by a total factor of over 60. Instead, and in line with our original working hypotheaia, our data suggest that the tetrazalylthio substituent, similarly to the acetoxy and carbamoyloxy groups, plays a definite role in contributing to good antibacterial performance (Ia vs. 10f). These incongruities, and unexpected losses of activity occasionally found for a few entries I, may hint at a subtle interplay between intrinsic activity and stability factors, which in the 6-unsubstituted series might have become particularly severe. A peculiar source of instability **common** to all the thiomethylpenems might be envisaged in the possibility of equi-<sup>44</sup>libration into the **exo** form under the conditions of antimicrobial testing. Conversely, on such hypothesis one could surmise that the activity observed for thioalkylidenepenams III, IV be the resuit **of** a slight proportion of the **penem** form arismg from equilibration, if not already preaent in the sample. Table Iv does indeed show that the potency ratio within each penem-penam couple is rather <sup>45</sup>evenly distributed **accross** the whole spectrum, **even** when, as for 1110, - the penam form apparently exhibits a remarkable level of antimicrobial activity.

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## REFERENCES **AND** NOTES

- 1. J.R.E. Hoover, "Antibiotics Containing the Beta-Lactam Structure", eds. by A.L. Demain and N.A. Solomon. Springer-Verlag. New York. 1983, vol 11, pp. 197-202.
- 2. D.B. Boyd, J. Med. Chem., 1984, **2,** 63.
- <sup>4</sup>3. This last effect **was** ascribed to the possible simultaneous departure of the leaving group X and ß-lactam acylation at the active site.
- 4. Evidences contrary to a concerted mechanism of this type have been recently put forward: M.I. Page and P. Proctor, J. Am. Chem. Soc., 1984, 106, 3820.
- 5. G. Franceschi. M. Foglio, F. Arcamone, A. Sanfilippo, and G. Schioppacassi. J. Antibiotics. 1980, 33, 453.
- 6. I. Ernest. J. Gosteli, C.W. Greengrass, W. Holick, D.E. Jackman. H.R. Pfaendler, and R.B. Woodward, J. Am. Chem. Soc., 1978, **100,** 8214.
- 7. M. Alpegiani, A. Bedeschi, G. Franceschi, F. Giudici, G. Nannini, and E. Perrone, Gazz. Chim. 1. Ernest, J. Gostell,<br>Woodward, <u>J. Am. Chem.</u><br>M. Alpegiani, A. Bedesc<br><u>Ital</u>., 1984, 114, 319.<br>M. Alpegiani. A. Bedesc Ital., 1984, 114, 319.<br>8. M. Alpegiani, A. Bedeschi, M. Foglio, and E. Perrone, <u>Gazz. Chim. Ital</u>., 1984, 114, 391.
- 
- 9. **A.** Yoahida, T. Hayashi, N. Takeda, S. Oida, and E. Ohki, Chem. Pharm. Bull. (Tokyo), 1981, 29, 2899.
- 10. 0. Mitsunobu, Synthesis, 1981. 1.
- 11. G. Franceschi. M. Foglia. M. Alpegiani, C. Battistini, A. Bedeschi, E. **Perrane,** F. Zarini. F. Arcamone, C. Della Bruna, and A. Sanfilippo, <u>J. Antibiotics</u>, 1983, 36, 938.<br>Arcamone, C. Della Bruna, and A. Sanfilippo, <u>J. Antibiotics</u>, 1983, 36, 938.<br>Although positive budgelined on silice sel plates, this compound s
- 12. Although readily hydrolyzed on silica gel plates, this compound could be isolated in 55% yield after flash-chromatography purification.
- 13. The final penems I, II, unless otherwise stated, were isolated as their Na or K salts and purified by reverse-phase chromatography (Merck LiChroprep RP-18; H<sub>2</sub>O or H<sub>2</sub>O-MeCN as eluants).
- 14. P.D. Jeffrey and S.W. McCombie, J. Org. Chem., 1982, **z,** 587.
- 7 15. The pNB esters of 2-heterocyclylthiomethyl penems cannot be cleaved by catalytic hydrogenation: however, we have found out in later experiments that vigorous stirring with iron powder in a biphase THF/conc. aq. NHC1 system enables the isolation of compounds I<sub>i</sub>in moderate yields.
- 16. **A.** Sanfilippo, C. Della **Bruna,** D. Jab&. E. Morvillo. *G.* Schioppacassi. G. Franceschi. F. Arca**mone,** C. Battistini, M. Foglio, and F. Zarini. J. Antibiotics, 1982, **35,** 1248.
- 17. S. Tsushima, M. Sendai, M. Shiraishi, M. Kato, N. Matsumoto, K. Naito, and M. Numata, Chem. Pharm. Bull. (Tokyo), 1979, 27, 696.
- 18. M. Alpegiani, A. Bedeschl, and E. Perrone, to be published.
- 19. Our results along this approach duplicate what reported by S. Uyeo at the Third Symposium on the Chemistry of ß-Lactam Antibiotics, Cambridge (UK) 1984.
- 20. E. **Perrone,** M. Alpegiani, A. Bedeschi, F. Giudici, M. Foglio, and G. Franceschi. Tetrahedron Lett., 1983, 24, 3283.
- 21. R.P. Volante, Tetrahedron Lett., 1981, 22, 3119.
- 22. The Mitsunobu reaction is not in common use for the synthesis of thioethers. For an isolated example, see H. Loibner and E. Zbiral, Helv. Chim. Acta, 1976, 59, 2100.
- 23. Iminophosphoranes have been previously obtained from carboxamides, sulphonamides, carbamates and nitroanilines, sometimes under THF reflux conditions;<sup>24</sup> to our knowledge this condensation has never been reported to occur on a heterocyclic amine.
- 24. H.J. Niclas and D. Martin, Tetrahedron Lett., 1978, 4031.
- 25. M. Pesson and M. Antoine, Bull. Soc. Chim. France, 1970, 1599.
- 26. R.B. Woodward, "Penicillin Fifty Years after Fleming", The Royal Society, London 1980, p. 245.
- 27. M. Foglio, G. Franceschi, G. Serra Errante, M. Ballabio, and F. Arcamone, Heterocycles, 1981, 15 785.
- 28. K. Hirai, Y. Iwano, and K. Fujimoto, Heterocycles, 1982, 17, 201.
- 29. P.C. Cherry, D.N. Evans, C.E. Newall, and N.S. watson, Tetrahedron Lett.. 1980, **2&** 5561.
- 30. It is worth noting the virtual coincidence of **J** all (cis) and J all (trans) between the vinyl and C-3 protons. For a somewhat different situation in related compounds, **see** ref. 27.
- 31. Compare this value with the data reported for  $E$  and  $Z$  isomers of clavulanic acid derivatives and, on the other hand, with the far greater difference ( $\Delta \delta \simeq 0.8$  ppm) observed between known 3R and 3S alkylidenepenam structures possessing a single (Z) alkene geometry.<sup>29,32</sup>
- 32. N.F. **Osborne,** J. Chem. Soc., Perkin Trans 1, 1982, 1429.
- 33. **We** regard this product as the result of a thialatian at the active C-3 methine of the **exo** form.
- 34. T. Hayashi, A. Yoshida, N. Takeda, S. Oida, S. Sugauara, and E. Ohki, Chem. **Pharm.** Bull. (Tokyo), 1981, 29, 3158.
- 35. S.R. Norrby and M. Johnnson, Antimicrob. Agents Chemother., 1985, 27, 128.
- 36. M. Foglio, C. Battistini, F. Zarini, C. Scarafile, and G. Franceschi, Heterocycles, 1983, 20,
- $, 1491.$
- 37. Selective cleavage of TBDPS vs. TBDMS ether (TBAF-3H<sub>2</sub>O; THF-HOAc, 30 min) in compound 19a is noteworthy, and should be related to the presence of the R-carbonyl group.
- 38. Every attempt to activate the acids  $17e$ ,  $17h$  through their mixed anhydrides with ethyl chlorocarbonate resulted in the formation of the corresponding ethyl ester. Mixed carboxylic-carbonic anhydrides usually require **some** particular catalysis (dimethylaminopyridine) in order to callapse into the esters: S. Kim, Y.C. Kim, and J.I. Lee. Tetrahedron Lett., 1982, 24, 3365.
- 39. Low yields in thioacid  $17d$  were mainly due to competing desilylation.
- 40. Desilylation under the usual conditions (TBAF) predominantly gave the **en0** form 23d. -
- 41. Direct heating of 21f gave penem 22f but, different from the TBDMS analogue 22c, this silyl ether was unaffected by acidic conditions compatible with the penem structure.
- 42. For a more detailed account on the antibacterial activity of heterocyclylthiomethylpenems, see G. Francesch~. M. Alpegiani. A. Bedeschi. M. Foglio, E. **Perrone,** G. Meinardi, S. **Grasso.** and I. de Carneri. J. Antibiotics,l984. *37.* 685.
- 43. J. Banville, P. Belleau, P. Dextraze, J.L. Douglas, F. Leitner, A. Martel. M. Menard. R. Salntonge, and Y. ueda. "syntheses and Structure-Activity Relationships in the **penem** Series". Presented at the 182nd ACS National Meeting, New York, 1981.
- 44. Analysis of additional "2-CH<sub>2</sub>X" penem structures wherein the double bond is locked in the endo position will be the subject of a forthcoming paper.
- 45. A low order of intrinsic activity, limited to Gram-positive species, might be proper of 29.32 alkylidenepenarns.

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