Concerning the Relative Non-Toxicity of Silacrown Ionophores

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PANNELL, K. H, C. K LA NEAVE, E RICO AND B ARKLES *Concerning the relative non-toxicity of silacrown tonophores* PHARMACOL BIOCHEM BEHAV 21: Suppl 1, 77-80, 1984 -Silacrowns, ionophoric materials in which an ethylene bridge of a normal crown ether has been replaced by a R_2S_1 group, facilitate ion transport across liquid membranes A major aspect of their properties is their nontoxicity when compared to these crown ethers. This is probably due to their ready hydrolysis to the corresponding polyethylene glycols The glycols also exhibit ion transport properties, even with an efficiency close to the silacrowns but the silacrowns stay intact long enough to partition into the hydrophobic phase of a hquid membrane whereas their hydrolysis products preferentially partition into the aqueous phases Ready removal from the hydrophobic phase of a membrane will reduce the long term transporting effectiveness of the glycols Thus, in vivo hydrolysis of the silacrowns will result in the loss of transmembrane carriers Similar removal will not occur for normal crown ionophores which are both hydrolytically stable and partition into the hydrophobic phase, and remain there for long time periods

THERE has been a considerable recent interest in the use of synthetic ionophores with respect to elucidating the various factors that effect the efficiency and selectivity of ion transport across real and artificial membranes by ionophonc materials [3, 4, 5, 7, 10] A major class of such synthetic compounds are the macrocychc polyethers, the so-called crown ethers. A major drawback to the large scale use of such compounds, in both industrial and medicinal environments is their tendency to have harmful physiological properties. For example, several of the most useful crown ethers are toxic, as are various naturally occurring antibiotic lonophores, e.g, monensm. Presumably the abdity of these species to slowly leak K^+ out of, and Na^+ into, the cell is responsible for this characteristic

Recently a series of new lonophores based partially upon organosdicon chemistry have been reported [1] In these compounds an ethylene bridge of a regular crown ether has been replaced by a disubstituted silyl group as illustrated in Fig. 1. Such substitution was expected to have several effects that may change the ion complexing and transporting ability of the resultant crown. For example, (a) size reduction of the central cavity, (b) change in the basicity of the crown due to the siloxy 0 atoms, and (c) reduction in the hydrolytic stability due to lability of siloxyethers.

We have reported that the efficiency of alkah metal ion binding and transporting of the silacrowns is reduced compared to the normal crown analogue and that the selectivity patterns are also changed. For example, whereas a normal crown with six oxygen atoms will usually be optimum for $K⁺$, in the case of the silacrowns the seven oxygen atom ring is optimum [9] Both of these properties seem to result from

the presence of the two sdoxy oxygen atoms Such atoms are less basic than normal ethereal oxygens [2], thus the binding of the sdacrowns to alkali metals will be less than the analogous non-substituted crown leading to less efficient transporting properties [7]. This feature will also create the need for a compensating O atom within the macrocycle nng as observed for optimum binding. Finally it has been demonstrated that the silacrowns are significantly less toxic than their unsubstituted analogues [1].

It is the purpose of this communication to report on the hydrolytic stabihty of the sdacrown lonophores, and suggest a relationship between this and their physiological properties. The results indicate that judicious use of organosilicon chemistry may lead to the synthesis of very useful medicinal compounds.

METHOD

The silacrowns were synthesized at Petrarch Systems as previously described [1], and used m this laboratory without further treatment.

Rates of potassium picrate transport were obtained using the apparatus and procedures described elsewhere [7] Concentrations of the ionophores in the hydrophobic phase and potassium picrate in the "in" aqueous phase were 10^{-2} m/l unless otherwise stated, and the temperature of the apparatus was mamtaaned at 20°C. The "in" and "out" aqueous phases (10 ml) were buffered using TRISMA systems at pH=7, and the chloroform solvent for the hydrophobic phase (25 ml) was preequilibrated with buffered aqueous solutions for 24 hr prior to a transport run

FIG 1 Substitution of the ethylene bridge of a regular crown ether by a silyl group Radical (R) represents phenyl, methyl, vinyl groups $n=1$ —Sila-11-Crown-4, 2 —Sila-14-Crown-5, 3—Sila-17-Crown-6, 4-Sila-20-Crown-7

In all experiments aliquots of the "out" aqueous phase were taken approximately every 20 m and the concentration of the picrate amon determined spectrophotometrically using the absorbance of the band at 357 nm. In all cases a plot of A versus t was hnear over the time periods of our experiments, 80-200 m. From such slopes the rate of transport was determined. These rates are initial rates since we are transportlng down a concentration gradient The final concentration of the potassium salts in the "out" phase is in the range 10^{-4} m/l, thus no problems are encountered by a close approach to the equilibrium condition

In experiments to determine the partitioning of the mnophores, both silacrowns and polyethylene glycols, a normal transport run was set up, *minus any salt tn the "m" aqueous phase* The set up was allowed to stir for periods of between 20 and 60 min. At this time the two aqueous phases were withdrawn and replaced with a salt solution in the "in" phase, and a buffered solution in the "out" phase and a normal $K⁺$ transport run was commenced. If partitioning of the lonophore into the aqueous phase has taken place, the removal of these phases wdl cause a loss of tonophore, hence reduction in transport. Related experiments were performed with tetraethylene glycol and the unsubstituted crown ether, 15-C-5

Studies on the hydrolysis of the sdacrowns were performed in the following manner An equal amount by volume of sila-14-C-5 and an isotonic saline solution ($pH=7.4$, 0 1%) NaC1, 0 03% gelatin) were permitted to react over a 24 hr period at 23°C. Periodically an allqot was taken and exam-

TABLE 1 TRANSPORT OF POTASSIUM PICRATE, (M/1 MIN) 108

Carrier	Rate	Ratio Crown/Glycol	
Sila-20-crown-7	400		
Hexaethyleneglycol	70	57	
Sila-14-crown-5	19	95	
Tetraethyleneglycol			

TABLE 2 HYDROLYSIS OF SILA-14-CROWN-5 IN ISOTONIC SALINE SOLUTION, pH 7 4*

Time (m)	$%$ silacrown	% tetraethylene glycol†
0	100	
	90	
25	44	34
66	37	38

*Analysis by liquid chromatographic analysis using a 500A styragel column, at 30°C

tWhile tetraethylene glycol is the ultimate hydrolysis product, an intermediate product is readily detected, probably $Me₂Si(OH)$ $(OCH₂CH₄)₄OH.$ If the ratio of silacrown to isotonic saline solution is increased, e g , to 5 1, then formation of the intermediate is the predominant initial reaction, with much less glycol formation Conversely, if the ratio is reduced much less intermediate is found at any time, and glycol formation is dramatically enhanced

lned by liqmd chromatography using a 500 A styragel column A measure of the hydrolysis was the relative amount of the parent sllacrown remaining at each measurement

RESULTS

The results of the normal transport runs for potassium picrate using the silacrowns and their corresponding polyethylene glycol hydrolysis products are presented in Table 1. These data clearly indicate that both the macrocycllc polyethers and their ring opened counterparts are effective at faclhtatmg transport Furthermore, the data indicate that the glycols are not grossly inferior to the sllacrowns in this respect since the ratios of [transport using crowns/transport usmg glycols] are small in these particular experiments, and never exceed an order of magmtude. The kinetic results (Table 2) of the hydrolysis of the silacrowns in saline solution clearly demonstrate the hydrolytic instability of the lonophores over a relatively short time period (24 hr), and confirmed the formation of the corresponding polyethylene glycols. This behaviour is expected for the S₁-O-C linkage [2]

The graphic displays of the results of the partitioning experiments for sila-14-crown-5, and hexaethylene glycol are presented in Figs. 2 and 3 They clearly indicate the difference in partitioning between the hydrophobic phase and water for the silacrowns as compared to the glycol hydrolysis products. Over the time periods in which preequilibrium between the hydrophoblc ionophore solutions and buffered aqueous

FIG 2 Plot of absorbance in "out" arm versus time for the Sila-14-Crown-5 \blacksquare -No preequilibrium, ∇ -15 m preequilibrium, O -30 m preequilibrium, \square -60 m preequilibrium

solutions are occurring, the silacrowns do not partition into the aqueous phase to any detectable amount, based upon transport rates Exactly similar results were obtamed using the unsubstituted crown ether, 15-C-S, indicating that such partitioning is not solely the result of the presence of the well established hydrophobic organosilicon moeity. The glycols behave m a different manner, partitioning significantly, hence being lost upon removal of the two "in" and "out" aqueous phases. The extent of partitioning will depend upon their concentrations in the hydrophobic phase and the volume of the aqueous phases. In the present set of experiments, for the hexaethylene glycol, complete equilibration of the glycol between chloroform and water occurs after approximately 30 m, Fig. 3. No attempt has been made m this work to determine the rate of such partitioning

DISCUSSION

We have demonstrated that the silacrown lonophores are susceptible to hydrolysis when treated with salme solution at pH 7.4, to form the corresponding polyethylene glycols. The rate at which this occurs is fast enough to suggest that in vivo the silacrowns will have a limited lifetime, prior to formation of the corresponding glycohc compound.

The comparative rate study between the sllacrowns and their glycol counterparts has illustrated that there is not a great deal of difference between their efficiencies as lonophores in the type of expernnent that we have performed. It is clear, and has been demonstrated by X-ray crystal structure analysis [6], that the glycols are capable of sequestering alkali metals and providing a hydrophilic environment for the cations, while at the same time presenting

FIG 3 Plot of absorbance in "out" arm versus time for hexaethylene glycol A--No preequilibrium, B-15 m preequilibrium, C —30 m and 60 m preequilibrium

an external hydrophobic shield needed for entry into, and passage through, a hydrophoblc region. If the toxicity of the various ionophoric substances is due to their ability to leak alkali metals across membranes why then are the silacrowns conspicuously non-toxic when compared to the normal crown ethers?

The ability of the crown ether lonophores to transport metals across membranes in vivo is predicated upon their ability to partition into the hydrophobic regions of various membranes, where they will act as carriers. This function will compete with, and disturb, the natural transport processes of the particular system with which they are involved. It is this function which presumably gives rise to their undesirable physiological properties. Since most synthetic lonophores have great thermodynamic, kinetic, and hydrolytic stability, once they have entered into the membrane hydrophoblc phase they will tend to stay there for long periods of time, permitting cation leakage, hence their toxic nature For the sllacrowns which also partition into the hydrophobic phase, there is a sufficient amount of water present to permit the slow hydrolysis to the glycohc materials. As we have demonstrated, these will rapidly partition into the aqueous phase and be removed by normal excretion processes. The result is that lonophonc materials will not remain in the hydrophoblc regions of membranes longer than the rate of hydrolysis of the silacrowns permit, and thus cation leakage is held to a minimum

We are currently investigating the rate of silacrown hydrolysis in an environment such as a membrane hydrophoblc phase with limited concentrations of water

An important implication stems from the results presented in this report. It has been reported that the naturally

occurring ionophore X-537A, which is selective for calcium binding and transport, has been used as an experimental drug to change hemodynamic responses of dogs undergoing open heart surgery [8] It is a long term goal of our research to synthesize silicon based ionophores which have the high selectivity and efficiency of cation transport noted for the naturally occurring species. By building into the structures hydrolytically labile bonds hydrolysis of the ionophore over short time periods will occur to yield readily expelled fragments, and non-toxic short term drugs

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