1 2	Kinetico-mechanistic studies of substitution reactions on cross-bridged cyclen CoIII complexes with nucleosides and nucleotides†
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- 29 Kinetico-mechanistic studies on the substitution reactivity of the $[Co{(\mu-ET)cyclen}(H2O)2]3+$
- 30 complex cation at pH values within the 6.0–7.0 range with biologically significant ligands have been
- 31 carried out. The substitution processes have been found to occur exclusively on the mono-
- 32 hydroxobridged [(Co{(μ -ET)cyclen}(H2O))2(μ -OH)]5+ species formed after equilibration of the cobalt
- 33 complex in the relevant medium. The studies conducted on the substitution of the aqua/hydroxo ligands
- 34 of this dinuclear species are indicative of a dominant role of outer-sphere complexation, involving
- 35 hydrogen-bonding interactions. The values of the outer-sphere complex formation equilibrium constant
- 36 are in line with the intervention of both the exiting aqua ligands and the NH groups at the encapsulating 37 $\{(\mu-ET)cyclen\}$ ligand. These complexes result in the preferential formation of O- or N-bonded
- nucleotides depending on the structure of the base moiety of the ligand. Even the entry of the different
- donor bonded nucleotides is hampered by the hydrogen-bonding interaction with the dangling moiety of
- 40 an already coordinated ligand. In general the overall substitution processes occur at a faster rate than
- 41 those published for the fully alkylated encapsulating $\{(Me)2(\mu-ET)cyclen\}$ ligand derivative, as
- 42 expected for the still available base-catalysing NH groups in the $\{(\mu-ET)cyclen\}$ ligand.

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45 INTRODUCTION

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47 As stated on many occasions, the use of coordination compounds to study possible modification in biologically significant molecules is not new, nevertheless any new information that can be extracted 48 from their simple reactivity should not be underestimated.1,2 Besides obvious thermodynamic 49 requirements and no leaching of the metal centre, the need for the processes to occur at a rate that allows 50 a controlled reaction, should also be taken into consideration when designing systems able to act in 51 52 biological systems. That is, the solvolysis or substitution processes that involve the metal centres have to be relatively slow in order to ascertain the maintenance of the active molecule, or the interaction of the 53 complex with the expected target.3 Nevertheless, the use of complexes that behave as dead-end species 54 55 for kinetic or thermodynamic reasons should also be avoided. A significant amount of literature has been lately appearing dealing with the speciation, hydrolysis, complexation and polymerization of a 56 57 number of "biologically" active centres, underlining the key role of simple substitution processes 58 actuating on biologically relevant coordination complexes.4-7 Clearly a rationalization of the solution 59 behaviour of metal complexes under conditions similar to those in biological media is fully desirable, including stability in plasma studies.8 Mimicking in vitro the conditions of biological systems is 60 61 extremely difficult, if possible at all, so that some simplifications have to be made. In this respect, hydrogen bonding and stabilization of supramolecular interactions have dramatic consequences in many 62 processes expected to be rather simple.9 Another important aspect relates to the effect of temperature on 63 64 the activity of these coordination compounds, 10 which underlines the importance of determining 65 activation parameters for these simple substitution reactivities, a point normally not considered. 66 Although the anticancer activity of cis-platinum still remains as the most important landmark in 67 medicinal inorganic chemistry, the importance of other metal complexes should not be underestimated.11 The last few decades have seen an increasing number of reports showing metal 68 complexes with promising medical applications and, for example, ruthenium complexes are now one of 69 70 the most important groups of compounds with antitumor properties.12-17 CoIII complexes with an inert 71 tetradentate skeleton and two reactive positions in cis are obviously interesting as they could represent a cheaper and less toxic alternative to currently used compounds.18 Even some Co alkyne complexes 72 73 have shown promising activity associated with their capability to target specific enzymes.19,20 The use 74 of complexes of type cis-[Co(N)4(H2O)2]3+, with (N)4 being tren, cyclen or fully alkylated {(Me)2(µ-75 ET)cyclen} has already been explored by us9,21,22 by studying their substitution processes with some nucleosides and nucleotides at physiological pH. In this respect there has been an important increase in 76 the use of cross-bridged cyclen and cyclam ligands, both for their promising properties and the 77

robustness of their structure containing five- and six-membered macrocyclic full encapsulation.23–25

79 Although for the tren and cyclen complexes the existence of fast base-conjugate processes dominate

their reactivity,26–29 the use of the fully substituted $[Co{(Me)2(\mu-ET)cyclen}(H2O)2]3+$ produces a definite increase in the inertness of the derivatives, due to the absence of acidic NH groups.27 In this

respect, the formation and cleavage of hydroxo-bridged dimeric units, kineticomechanistically detected

some time ago, 30, 31 has been found to be a keystone. The prevalence of reasonably reactive dinuclear

84 mono-hydroxo-bridged species is directly related to the presence of these NH groups, only dead-end

 $(N)4Co(\mu-OH)2Co(N)4$ cores detected at pH > 7.2 for the fully alkylated [Co{(Me)2(μ -

86 ET)cyclen(H2O)2]3+ material.

87 In view of these facts, the study of the complex with the tuned, partially substituted but conformation-

rigid, $\{(\mu-ET)-cyclen\}$ (Scheme 1, top) cyclen-based ligand has been pursued. By the use of this ligand,

89 base-conjugate accelerated substitution processes would still be active at pH values normally higher than

90 neutrality, once the extremely acidic equatorial NH groups in cyclen have been substituted.29

91 Furthermore, the presence of acidic hydrogens attached to the axial nitrogen donors should also allow

- some of the interactions observed for the [Co(cyclen)(H2O)2]3+ complex9 with Good's buffer media,32
- as well as in ZnII complexes with nucleotides.33 In this report we present the study of the spontaneous

- solution chemistry of $[Co{(\mu-ET)cyclen}(H2O)2]3+$ in a pH range close to neutrality, as well as the
- 95 kinetico-mechanistic studies on its substitution reactivity with chloride, inorganic phosphate and the
- 96 nucleosides and nucleotides indicated in Scheme 1. The results collected agree with the reactivity of the
- 97 CoIII t2g 6 metal centre in dinuclear [{Co{(μ -ET})cyclen}-(H2O)}2(μ -OH)]5+ units. The processes are
- also slightly accelerated, with respect to those observed for the fully substituted $\{(Me)2(\mu-ET)cyclen\}$
- analogue, by the actuation of a base-conjugate pathway. Reactivity stops at pH > 7.5, where the
- 100 prevalence of the bis-hydroxo dimeric form of the ion occurs. The data collected are also indicative of
- 101 important outer-sphere interactions between the donors on the nucleobase moieties and both the
- 102 remaining NH axial groups of the encapsulating ligand, and the aqua ligands on the CoIII coordination
- 103 centre.

105 RESULTS AND DISCUSSION

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107 Solution behaviour of [Co{(µ-ET)cyclen}(H2O)2]3+ in the 5.5–7.5 pH range

- 108 The diaquo $[Co{(\mu-ET)cyclen}(H2O)2]3+$ complex has been obtained for the first time in this work as a 109 triflate salt, and XRD analyses of the compound has also been carried out (Fig. 1). Distances and angles
- 110 determined for the cation complex do not show any significant difference with those from the cyclen34 (110)
 - and $\{(Me)2(\mu-ET)cyclen\}$ 18 related complexes. The values of the two pKas for this species have been determined, both by potentiometric and spectrophotometric NaOH titrations at I = 1.0 (NaClO4), as 5.1
 - and 7.4 at 25 °C, as indicated in the Experimental section. With these values in hand, the species present
 - in the pH margin of our kinetic studies are $[Co{(\mu-ET)cyclen}(H2O)(OH)]^2+$ and $[Co{(\mu-ET)cyclen}(H2O)(OH)]^2+$
 - 115 ET)cyclen}(OH)2]+. Determination of these pKa values using different pH equilibration times (i.e. 10–
 - 116 70 seconds) was also carried out with the same outcome, thus indicating that the processes being
 - measured correspond effectively to the deprotonation equilibria with no significant intervention of
 - secondary processes. Once the acid/base prevalent species at physiological pH was established, the
 - 119 possible polymerization processes occurring due to the generation of bridging hydroxo ligands on
 - increasing the pH was pursued.9,22,28,35 Studies conducted on $(5-10) \times 10-4$ M solutions of the
 - 121 [Co{(μ -ET)cyclen}(H2O)2]3+ complex, in non-buffered final pH = 7.5, show a set of two step changes
 - 122 in the UV-Vis spectrum on the 30 minute time-scale. By comparison with the previous solution 122 $\frac{1}{2}$ $\frac{$
 - 123 chemistry of the parent [Co(cyclen)(H2O)2]3+ and the fully alkylated [Co{(Me)2(μ -ET)-
 - 124 cyclen}(H2O)2]3+ complexes,9,22 the reactions have been associated with the sequential formation of
 - 125 mono- and bis-hydroxobridged, $[(Co{(\mu-ET)cyclen})2(\mu-OH)2]4+$, complexes.
 - By the use of MES and HEPES buffers, a pH screening of this spontaneous solution reactivity has been
 - 127 conducted. In all cases the mentioned UV-Vis spectral changes are reproducible (Fig. S1[†]), and show
 - the two-step sequence indicated above. The time-scale of these processes (10 plus 70 minutes at 17 °C) is also being the set of the
 - is clearly intermediate between those observed for the parent cyclen9 and those found for the fully $(0.42)^2(0.57)$ and $(0.42)^2(0.57)$ and $(0.42)^2(0.57)$
 - substituted {(Me)2(μ-ET)-cyclen}22 analogous derivatives. This trend is in line with the residual
 presence of two slightly acidic NH axial groups attached to the CoIII centre in the present compound,
 - still capable of induced base-catalysed substitution reactivity.29,36 At the same time-scale, for the
 - 133 parent cyclen derivative, the presence of very acidic equatorial NH groups,29 induce fast polymerization
 - reactions.9 As found for the previously studied related systems, the absorbance changes increase
 - 135 significantly with increasing pH, but can be reversed in a fast process by addition of HClO4 to acidic
 - pH. At pH > 7.5 the changes associated with the second process become very important and are related
 - to the lack of reactivity observed with the variety of ligands studied (see the following section).
 - Summarising, the behaviour of the $[Co{(\mu-ET)cyclen}-(H2O)2]3+$ system has intermediate features
 - between those of the parent cyclen9 and $\{(Me)2(\mu-ET)cyclen\}22$ derivatives. The polymerisation
 - equilibria indicated in Scheme 2 are at pH values higher than 7.5 (after 30 minutes at room temperature,
 - 141 see below) significantly displaced to the formation of a dimeric dead-end bis-hydroxo [(Co{(μ -
 - 142 ET)cyclen})2(μ -OH)2]4+ complex. At lower pH values (within the 6.0–7.0 range) the mono-
 - 143 hydroxobridged dinuclear species is prevalent on equilibration for a few minutes, and reactivity is,
 - 144 consequently, expected from the aqua ligands.9,22
 - 145

Substitution reactions on [Co{(µ-ET)cyclen}(H2O)2]3+ in the 6.0–7.5 pH range by chloride, phosphate, cytidine, thymidine, uridine, 5'-cytidinemonophosphate, 5'-thymidinemonophosphate

148 and 5'-uridinemonophosphate

- 149 After the knowledge of the solution nature of the equilibrated species of the $[Co{(\mu-$
- 150 ET)cyclen}(H2O)2]3+ complex in the 6.0–7.5 pH range, and in view of some studies carried out on this

- 151 core in biologically relevant processes, 18,37 the reactivity at physiological pH of the complex was
- 152 pursued. For the parent cyclen and $\{(Me)2(\mu-ET)cyclen\}$ complexes at these pHs,9,22 no reactivity with
- 153 chloride had been observed, which is relevant in view of the chemistry of cis-platinum at different
- pCl.38–41 For the present complex, the substitution of the aqua/hydroxo ligands at pH values between
- 6.0 and 7.5 and at [Cl-] = 0.05-0.075 M is also not observed after 24 hours at 40 °C.
- 156 Phosphate. As a follow up, the reactivity of the complex with simple inorganic phosphates was studied 157 as a model for the substitution processes occurring with nucleotides. Furthermore, the process has a 158 perfect spectroscopic NMR handle21,42 for the establishment of the nature of the reacting and final 159 complexes. Monitoring the spectral changes on non-equilibrated, freshly prepared, solutions of compound $[Co{(\mu-ET)-cyclen}(H2O)2]3+$, with phosphate leads to a complex sequence where the 160 fastest changes were equivalent to those observed for solutions not-containing the ligand. Thus, by using 161 the methodology indicated in the Experimental section,9,21,22 the time-resolved UV-Vis spectral 162 changes occurring on pre-equilibrated (30 minutes at room temperature) samples of $[Co{(\mu-ET)}-$ 163 164 cyclen}(H2O)2]3+ at the relevant pHs were monitored instead. The changes observed (Fig. 2a) in the UV-Vis spectrum, can be fitted to a two-step reaction sequence with rather similar rate constants that 165 166 were obtained as described in the Experimental section.43,44 While a definite limiting dependence, i.e. $kobs1 = KOS \times k1[phosphate]/(1 + KOS[phosphate])$, on the total concentration of phosphate is evident 167 for one of the steps observed, no dependence on concentration is obtained for the other step in the 168 sequence (Fig. 2b). Clearly the limiting first-order rate constant obtained from these plots corresponds to 169 170 the coordination of a phosphate anion on a precursor outersphere encounter complex (as already been 171 established for other phosphorous oxoanion substitution reactions).45-47 Consequently, the other 172 reaction should correspond to a consecutive chelation, or bridging, of the ligand after its coordination.
- 173 Table 1 shows the relevant kinetic data for these processes.
- 174 For the concentration dependent (k1) path, the limiting behaviour agrees with that observed for the
- parent cyclen complex, but is contrary to that for the $\{(Me)2(\mu-ET)cyclen\}$ analogue.9,22 Clearly, as
- 176 found in other systems,33 the available and well-oriented NH groups in the macrocyclic ligand promote
- such outer-sphere association complexes with the entering anion (avKOS = 120 M-1 in this case). As
- 178 for the outcome of the concentration independent (k2) path, 31P NMR spectroscopy of the final reaction
- 179 mixtures indicated the presence of a phosphato ligand with either a $\eta 2$ or μ geometrical arrangement (20
- 180 ppm downfield from the signal of the free anion).42 Experiments carried out with a [Co] : [P]total = 1:5
- ratio produced a final [P]total : [P]20 ppm = 4.5 : 0.5, thus indicating that a simple [(Co{(μ -ET)cyclen})2(μ -OOPO2)(μ -OH)]2+ complex is formed after the reaction (Scheme 3, top). The same
- arrangement has been observed for the parent unsubstituted cyclen derivative,9 which is in line with that
- indicated in Scheme 2. Finally it is interesting to note, as seen both in Fig. 2b and Table 1, that, although
- some pH trends might be present in the rate constants determined, these are not significant; the reactions
- seem to be pH independent in the narrow range studied (6.0–7.0). In view of the reactivity observed,
- 187 completely parallel to that for the parent cyclen derivative,9 the thermal and pressure activation
- 188 parameters for this system have not been determined.
- 189 Cytidine. The study of substitution reactions on the above mentioned CoIII complex with biologically relevant ligands with nitrogen donors, i.e. nucleosides or nucleobases, was further intended (after the 190 equilibration indicated in the previous section). No changes were observed at $pH \ge 7.5$, in good 191 agreement with the formation of the dead-end bis-hydroxobridged dimers (Scheme 2), and the reaction 192 193 was only studied in the 6.0–7.0 pH range, where the $[(Co{(\mu-ET)cyclen}(H2O))2(\mu-OH)]5+$ species is prevalent in the medium. By using the standard software, 43, 44 these changes were associated with a 194 195 twostep sequential process. Water Presat proton NMR experiments on the reacting solutions established 196 the nature of the two species appearing during the process. The spectrum collected after 1 hour of 197 reaction at 40 °C at pH = 6.1 shows, apart from the intense doublet at 7.8 ppm corresponding to the
- para-NH proton of the ring of the free cytidine, a signal at 7.7 ppm. After further 24 hours under the
- same reaction conditions, the signal at 7.7 ppm increases its intensity and a new signal appears at 7.6

- 200 ppm. These data agree with the initial formation of a mono-cytidine complex (7.7 ppm) that evolves to a
- 201 bis-cytidine species (7.6 ppm) with time (Fig. S2[†]) in an equilibrium overall process. Fig. S3[†] shows the
- trends observed for the two pseudo-first order rate constants with the concentration of cytidine; the 202 values derived at different pHs are shown in Table 1. From these plots it is clear that an equilibrium 203
- 204 condition is established for both substitution reactions (as indicated by the NMR data). The equilibrium
- constants (10-40 M-1 for K1 and 50-90 M-1 for K2) indicate a definite preference for the 205
- 206 bissubstituted [(Co{(μ -ET)cyclen}(cytidine))2(μ -OH)]5+ complex. The slight acceleration (see Table 1)
- of the rate on increasing the basicity of the medium is in good agreement with the residual operation of a 207
- 208 conjugate-base mechanism due to the existence of NH groups in the cobalt encapsulating ligand used.
- 209 Given the complexity of the data only the thermal activation parameters at pH = 6.5 have been
- determined. For ΔH_{\pm}^{\pm} , the values indicate in all cases a high degree of dissociativeness, as expected for 210
- CoIII complexes when conjugate-base mechanisms might be operative. The values for the activation 211
- 212 entropies are also positive, but less than expected for this type of activation, especially as k1 and k2 on
- rate constants are concerned. The inclusion of the outer-sphere association constants in the value used 213 for the determination of these parameters might be responsible for the lesser degree of dissociativeness
- 214
 - 215 observed.
 - 216 Thymidine. Initially, as for previous studies,22 thymidine (Scheme 1) was chosen due to the neutral
 - characteristics of its nitrogen donor (pKa = 9.8),48,49 which allows for a simplification of the system, 217
 - despite its anionic nature once coordinated to the CoIII centre.33 Fig. 3a shows the typical UV-Vis 218
- 219 spectral changes observed in the system and Fig. 3b shows the trends for the pseudo-first-order rate
- 220 constants obtained at different acidities for this substitution process with thymidine. While the slow step
- 221 (kobs2) shows a linear dependence on the concentration of the entering ligand, the fastest step (kobs1) 222 clearly shows a limiting behaviour22,35,50,51 on the thymidine concentration (as already found for the
- 223 phosphate substitution process, see above). As in previous studies, no dependence on the pH is
- 224 observed, and the limiting value of the fast step (k1 (s-1)) and the slope of the dependence of the second
- step (k2 (M-1 s-1)) were obtained from an average of the data at the same thymidine concentrations at 225
- different pHs. The relevant data extracted from these plots indicated are shown in Table 1. 226
- 227 Thus, as for cytidine, the substitution reaction of complex $[Co{(\mu-ET)cyclen}(H2O)2]3+$ with
- 228 thymidine at pH values between 6.0 and 7.0 corresponds to a process occurring on the mono-hydroxo
- bridged [(Co{(μ -ET)cyclen}(H2O))2(μ -OH)]5+ species prevalent in this medium (see Scheme 3, 229
- 230 bottom). The reaction produces the bis-substituted $[(Co{(\mu-ET)cyclen}(thymidine))2(\mu-OH)]3+$ species
- as the final complex via a clear dissociative activation mechanism for each sequential step. This is 231
- evidenced by the large values of ΔH_{\pm}^{\pm} (Table 1) of the same magnitude than those found for similar 232 233 dissociatively activated CoIII amine complexes.52–54 Interestingly, the values determined for ΔS_{\pm}^{\pm} are
- practically zero, and the volumes of activation are negative for the k1 process and positive for the k2 234
- step. This unexpected trend in the values of $\Delta S^{\ddagger}_{\ddagger}$, and their lack of correlation with $\Delta V^{\ddagger}_{\ddagger}$, has been 235
- 236 related to the existence of important solvent assisted hydrogen bonding interactions in the transition
- state of the substitution process.55–59 This is not surprising in this case, the nature of both the entering 237
- 238 and encapsulating ligands (see Scheme 1) should be prone to hydrogen bonding interactions. In this
- respect, the previously reported22 reaction of $[Co{(Me)2(\mu-ET)cyclen}(H2O)2]3+$ with 5'-TMP is a 239
- 240 clear example of this effect in this family of substitution processes.
- 241 In contraposition with that observed for the cytidine substitution, where no acidic protons are available
- 242 in the ligand, for the entry of the first thymidine nucleoside the reaction involves the limiting formation
- of an outer-sphere encounter precursor complex (KOS = 240 M-1, Fig. 3b and Table 1), as found for 243
- 244 other substitution reactions on CoIII amine systems,47,60 as well as on the related [Co{(Me)2(µ-
- T)cyclen}-(OH)(H2O)]2+ complex.22 The formation of this outer-sphere complex has been associated 245
- with the interaction of the proanionic {ONO} unit of the nucleoside (see Scheme 1) both with the 246
- 247 protons of the remaining aqua ligand in the CoIII complex at this pH and the NH groups in the ligand.33
- Effectively, for this nucleoside the value of KOS is an order of magnitude larger than for the not-248

- containing NH group analogue systems studied.22 As for the entry of the second thymidine ligand, the
- value of the outer-sphere association equilibrium constant has definitively decreased in a way that a
- simple linear dependence between the value of kobs2 and [thymidine] is obtained (Fig. 3b). The
- decrease of the positive charge of the CoIII complex, produced by the first substitution of H2O by
- thymidine (which involves its deprotonation to the thymidine $\{-H\}$ –anionic ligand)7,33 can easily
- explain this fact.
- 255 The fact that the water by thymidine substitution reactions are not affected by the pH in the narrow
- range studied is somehow surprising. A compensation effect between the increasing amounts of the
- more labile H2O/OH– CoIII species on decreasing the pH, and the viability of an accelerated
- 258 baseconjugated substitution on its increase, seems to result in the final outcome observed. Furthermore, 259 the protonation ambiguity effect on putative full hydroxo outer-sphere complexes can also explain this
- 260 fact for the first nucleobase entry.61
- **Uridine.** The substitution process with a very similar uridine nucleoside (Scheme 1) was also pursued and a completely parallel behaviour with respect to the substitution by thymidine (Fig. S4†) has been observed. The slightly higher values of the rate constants can be associated with its lower steric requirements (see Scheme 3). Furthermore, its higher (pKa = 9.3)62 acidity also weakens the outersphere association of the {ONO} unit of the nucleoside, as observed from the less pronounced curvature for the k1 versus [uridine] trend (Fig. S4a†). Given the similarity obtained with the thymidine studies above, as well as the expected character of the variations observed, no further studies related to pH
- 268 variation and activation parameters were conducted.
- 5'-CMP. Once the reactivity with phosphates and nucleosides was established, the substitution
 processes by nucleotides was pursued;9,22 the 5'-CMP nucleotide has been our first choice due to the
 similar anionic behaviour to phosphate anions, as well as for its non-pro-anionic nature on the
- nucleoside moiety. Effectively, as for the H2PO4-/HPO42- system the spectral changes indicate a two-
- step reactivity pattern at 40 °C within the 6.0–7.0 pH range, where the ligand is in the 5'-CMP-/5'-
- 274 CMP2- forms.63 The global fitting of these timeresolved changes43,44 produced a series of rate
- constants that are concentration dependent for the two consecutive steps (Fig. S5[†]). 31P NMR
- spectroscopy indicated that an initial species is formed showing a signal at 9.4 ppm downfield from that
- of the free ligand, followed by the appearance of another signal at 14.1 ppm also downfield from that ofthe free ligand. Both signals coexist in the final reaction mixture and correspond to mono-dentate
- 278 the free figand. Both signals coexist in the final feaction inixture and correspond to mono-definate 279 phosphato ligands involved in equilibrium (Fig. S6†). The full reaction scheme for this system, parallels
- what has been observed for the parent [Co(cyclen)-(H2O)2]3+ compound (Scheme 4, top).9 The fact
- that no induction period has been observed in the NMR experiments, and that the shift in the UV-Vis
- 282 occurs to lower energies, is also an indicative of the neat formation of O-bound 5'-CMP complex. The
- data collected in Table 1 indicate that, in contrast to the observed for the reaction with inorganic
- phosphate, the reaction rate increases with pH. The increasing presence of fully deprotonated 5'-CMP2– (pKa = 6.1) species, not so much relevant for inorganic phosphate (pKa = 7.21), seems to be
- 286 decisive.42,63
- 287 The absence of a measurable value of KOS for the substitution by 5'-CMP, when compared to that with
- inorganic phosphates, is in line with an intermediate behaviour from that observed for the reactions with
- $[Co(cyclen)(H2O)2]3+ and those with [Co{(Me)2(\mu-ET)cyclen}(H2O)2]3+.9,22 Clearly the presence of [Co(cyclen)(H2O)2]3+.9,22 Clearly the presence$
- 290 NH groups in the ligand structure (see Scheme 1) plays a determinant role in the outer-sphere
- association with the smaller H2PO4–/HPO42–, which is not possible with the more encumbered 5′-
- 292 CMP-/5'-CMP2- anion. Nevertheless, additional hydrogen bonding interactions with the solvent may
- also be a competing factor to account for the weaker interactions observed between the CoIII complex
- and the ligand in this case. From the data in Table 1 the values of K1 and K2 are found equivalent
- considering the methodological errors involved, their values are apparently also independent of the pH
- and within the 7–11 M–1 range.

- As for the thermal activation parameters shown in Table 1 for this system, it is interesting to note that,
- 298 while for the first entry of 5'-CMP the direct process has all the characteristics of a dissociatively
- activated reaction, for the entry of the second ligand a much higher degree of associativeness is
- 300 observed. The same is observed for the aquation reaction occurring on the same material, i.e. [(Co $\{(\mu \mu)\} \in \mathcal{A}_{n}\}$)]
- 301 ET)cyclen}(5'-CMP))(μ -OH)-(Co{(μ -ET)cyclen}(H2O))]3+ (Scheme 4). It is clear that, although a
- dissociatively activated substitution mechanism is expected for tetraamine CoIII complexes, especially
 when a conjugatebase process can be operative,28,36 outer-sphere interactions lead to a certain degree
- 303 when a conjugatebase process can be operative, 28,36 outer-sphere interactions lead to a certain degree 304 of association which also facilitates the reaction. It is clear that the nature of the complex plays a key
- role in the above mentioned interactions, especially taking into account the presence of two NH groups
- 306 in the macrocyclic ligand of the inert skeleton. In fact a recent report has appeared in reference to such
- 307 outer-sphere interactions as responsible for deeply tuning the electronics of rather simple complexes.64
- **5'-TMP.** The substitution reactions with the 5'-TMP nucleotide (Scheme 1) were studied to generalise the interesting substitution trends observed for the different phosphates with this CoIII complex.22 The
- 310 nature of this nucleotide allows the formation of anionic N- and O-bound nucleotide complexes. As for
- the previous systems in the present report, the timeresolved UV-Vis spectral changes can be fitted to a
- two-step sequence with the concentration and pH dependence characteristics indicated in Fig. 4. 31P
- 313 NMR experiments were conducted to establish the nature of the product of the two set of consecutive 214 magnetized at 40 °C and all = (5 the analysis of the 21D NM)
- 314 processes. After 30 minutes at 40 °C and pH = 6.5 the only signal appearing in the 31P NMR 245 processes later that of the first H = 1/2 Q = 1/2 h = 1/2
- corresponds to that of the free ligand (2.9 ppm), while after a further period of 10 hours a new signal at
 12.6 ppm appears, corresponding to a mono-dentate O-phosphate. The relative intensity of these 31P
- NMR signals indicate the entry of a single phosphate ligand per each CoIII 2 unit, thus indicating the
- validity of the implied reactive species indicated in Schemes 2 and 4 in this narrow pH range. It is thus
- 319 clear that the same behaviour observed for the fully alkylated derivative is operative:22 formation of
- 320 $[(Co{(\mu-ET)cyclen}(N-5'-TMP))(\mu-OH)(Co{(\mu-ET)-cyclen}(H2O))]4+$ that evolves to the final
- 321 $[(Co{(\mu-ET)cyclen}-(N-5'-TMP))(\mu-OH)(Co{(\mu-ET)cyclen}(O-5'-TMP))]2+$ species. On standing for
- longer periods, the final fully substituted species isomerises with the formation of $\{(Co\{(\mu Co\})\}$
- ET)cyclen}(O-5'-TMP))2(μ -OH)} units, as evidenced by a new signal on the 31P NMR spectrum
- appearing at 17.4 ppm (Scheme 4, bottom right). Table 1 shows the relevant kinetic and activation
- 325 parameters for the substitution processes studied.
- 326 Interestingly, the values for the entry of the first 5'-TMP ligand (N-bound) have a rather low value of
- Δ H[‡] and extremely negative activation entropy. It is clear that, even in this inherently dissociatively activated substitution process (CoIII t2g6), an important outer-sphere degree of associative activation
- must be present, also evidenced by the values determined for ΔV_{\pm}^{\pm} . This is in good agreement with the
- presence of an {ONO} unit in the ligand plus some NH groups in the inert skeleton of the complex.33
- 331 For the entry of the second (O-bound) 5'-TMP, this effect seems to be minimised, and a large value of
- 332 ΔH_{\pm}^{\pm} is determined together with values of ΔS_{\pm}^{\pm} and ΔV_{\pm}^{\pm} close to zero.
- As for the absence of pH dependence in the values of k1, previously reported data referring to proton
- ambiguity for these systems,61 as well as protonation of the outer-sphere complexes,65 can be held
- responsible for the facts.22 Nevertheless, taking into account the potential actuation of conjugatebase
- mechanisms, as well as the very narrow range of pH used in the study (6.2-7.0), the above assumptions
- are highly speculative (see Fig. 4).
- **5'-UMP.** The effect of the substitution of uridine for thymidine on the corresponding nucleotides was
- also studied. Fig. S7^{\dagger} shows the trend observed on [5'-UMP] of the two derived values of kobs at pH =
- 340 6.5. As for 5'-TMP the nature of the species being formed after each one of the sequential steps was
- assessed by 31P and Presat 1H NMR experiments (Fig. S8†). After a period of 1 h at 40 °C and pH = 6.8
- a signal appears at 12.4 ppm in the 31P NMR spectrum, indicating the monodentate O-5'-UMP
- 343 coordination to the cobalt centre. The parallel resat 1H NMR experiment indicates the presence of two
- doublets (7.8 and 7.9 ppm), the more intense at 7.9 ppm is assigned to the mono-N-5'-UMP. The signal
- of the 31P NMR spectrum indicates that the proton signal at 7.8 ppm must then correspond to a mono-

- 346 N-mono-O species as already established for the 5'-TMP derivatives. The rate constants derived from
- 347 the experiments are practically equivalent to that obtained for the 5'-TMP, indicating a similar nature of
- 348 the reaction. Nevertheless the corresponding value for KOS is much lower than for the thymidine
- 349 derivative, in line with the observed for the reaction with the parent nucleosides. Given the complex
- 350 nature of the secondary reactions observed, no pH, temperature or pressure dependence has been
- pursued for this system. 351
- 352

Comparison within the cyclen, {(µ-ET)cyclen} and {(Me)2(µ-ET)cyclen} series of complexes 353

- 354 In Scheme 5 a comparison of the structures of the series of CoIII aqua complexes with cyclen
- derivatives is indicated, as well as the final thermodynamically equilibrated form existing in solution at 355
- pHs close to the physiological pH.9,22 Successive substitution on the NH groups of the cyclen ligand 356
- produces a significant decrease in the relevance of the dimerization processes under biologically 357
- significant conditions; which is related to the expected tuning on conjugatebase pathways dominant for 358
- 359 the cyclen derivatives,29 relevant for the simple cross-bridged derivate, and irrelevant for the complexes of the fully alkylated derivative.22 These trends indicate a thermodynamic preference of the dimeric µ-
- 360
 - OH species.9 361
 - 362 A further very interesting tuning found for the series of studies, relates to differences in hydrogen
 - bonding interactions due to the presence of NH groups in the ligand. While for the fully alkylated 363
 - 364 compound hydrogen bonding interactions with thymidine seems to be solely taking place via
 - interactions with the hydroxo ligand on the monomeric CoIII centre, for the cross bridged ligand 365
 - compound here reported, the involvement of the axial NH group increases by an order of magnitude the 366
 - stability of such an interaction (Scheme 6, left). This is especially important considering that the extreme 367 368 dissociative substitution occurring on conjugate-base pathways is not too relevant for this complex (see
 - above), thus allowing for an unexpected association. 369
 - 370 Precisely this outer-sphere association is selective, and the formation of N-bound thymidine for the 5'-
 - TMP nucleotide can be observed despite the phosphate-bound derivative being thermodynamically 371
 - preferred as shown in Scheme 4. Moreover, a possible outer-sphere pairing interaction can be also held 372
 - responsible for the unobserved bis-N-bound thymidine derivative, as indicated in Scheme 6 (right). 373
 - 374

375 CONCLUSIONS

376

- 377 The substitution reactivity of the $[Co{(\mu-ET)cyclen}(H2O)2]3+complex cation at pH close to neutrality$
- is dominated by the formation of the dead-end bis-hydroxobridged $[(Co{(\mu-ET)-cyclen})2(\mu-OH)2]4+$ 378
- complexes at pH > 7.5, which parallels what has been observed for the parent cyclen derivative and that 379
- of the fully alkylated $[(Co{(Me)2(\mu-ET)cyclen})2(\mu-OH)2]4+$. At Ph values within the 6.0–7.0 range 380
- substitution reactivity occurs solely on the mono-hydroxobridged $[(Co{(\mu-ET)cyclen}-(H2O))2(\mu-ET)cyclen}]$ 381 382 OH)]5+ species, as observed for the cyclen analogous system, but differing from the fully alkylated
- derivative where this species is only residual. It is clear that the presence of NH groups in the axial 383
- coordination positions of the complex (Scheme 1) dominates this reactivity, whereas that of the highly 384
- 385 acidic NH in the equatorial positions of the [Co(cyclen)-(H2O)2]3+ complex is only responsible for the
- readiness and equilibrium position of the mono-hydroxobridged species formation reaction. 386
- 387 The kinetico-mechanistic studies conducted on the substitution of the aqua/hydroxo ligands of this
- 388 dinuclear species with biologically significant ligands are indicative of an important role of outer-sphere
- complexation. The appearance of limiting kinetics is rather general, and with values of KOS that are in 389
- line with the intervention of both the leaving aqua ligands and the residual NH groups in the 390
- 391 encapsulating $\{(\mu-ET)-cyclen\}$ ligand (Scheme 6). The formation of hydrogenbonded encounter
- complexes results in the preferential formation of O- or N-bonded nucleotides depending on the 392
- structure of the base moiety of the ligand. Furthermore, as observed for the reaction with mononuclear 393
- 394 $[Co{(Me)2(\mu-ET) cyclen}(H2O)-(OH)]2+$, the selective entry of the different donor bonded nucleotides 395
- can also be hampered by the hydrogenbonding interaction with the dangling moiety of the coordinated
- 396 ligand. As expected, the processes occur at a faster rate than for the fully alkylated encapsulating ligand 397 derivative, given the still available base-catalysing NH groups in $\{(\mu-ET)cyclen\}$.

399 EXPERIMENTAL

400

401 Compounds and procedures

- 402 The cobalt $[Co{(\mu-ET)cyclen}(H2O)2](CF3SO3)3$ complex has been prepared directly by
- 403 crystallisation of the already described $[Co{(\mu-ET)cyclen}(CF3SO3)2](CF3SO3)$ in a minimum amount
- 404 of slightly acidic (HCF3SO3) water.18 The complex was characterised by its elemental analyses (Calc.,
- 405 found %) for $[Co{(\mu-ET)cyclen}(H2O)2](CF3SO3)3 \cdot 1.5H2O: C: 20.3, 20.0; N: 7.3, 7.2; H: 3.8, 3.7; S:$
- 406 12.5, 12.6. 13C NMR (57.6, 62.8, 63.2 ppm) and UV-Vis spectra (λ max = 360 nm (110 M-1 cm-1);
- 407 $\lambda \max = 490 \operatorname{nm} (180 \operatorname{M}{-1} \operatorname{cm}{-1})$). XRD quality crystals were also obtained; Fig. 1 shows the molecular 408 drawing of the complex cation and Table 2 the corresponding crystal data and structure refinement. The
- 409 nucleosides and nucleotides used were commercially available and were used as received.
- 410 MES and HEPES solutions were prepared to a 0.4 M concentration at I = 1.0 (NaClO4) by weighing the
- desired amounts of the commercially available reactants. Final pH was adjusted with suitable HClO4 or
- 412 NaOH solutions.32 These stock solutions were used as the solvent for all the ligand solutions used in the
- study. As a standard procedure the pH of the samples at the desired temperature was monitored before
- and after randomly selected experiments; no significant differences were obtained in any case and the
- 415 procedure was thus considered valid under the conditions used.
- 416 13C and 31P NMR spectroscopy were carried out on Bruker 400-Crio instrument in H2O adjusted at the
- 417 desired pH, and with a D2O inset containing the corresponding reference, at the Unitat d'RMN from the
- 418 Centres Científics i Tecnològics de la Universitat de Barcelona (CCiTUB). The spectra were referenced
- externally to NaTMPS (13C) and H3PO4 (31P). 1H NMR spectra from the same aqueous solutions were
- recorded using a water Presat experiment on the same instrument. UV-Vis spectra were recorded using
 either a Cary 50 or a HP8453 instrument equipped with thermostated multicell transports. For the
- 421 cruter a Cary 50 or a r1r 6455 instrument equipped with thermostated multicell transports. For the
 422 reactions carried out at varying pressure the already described pillbox cell and pressurising systems66–
- 423 69 were used connected to a J&M TIDAS instrument. pH measurements were conducted on a Crison
- 424 instrument using either fast response or microsample glass combined electrodes. Time-resolved UV-Vis
- 425 spectra were recorded with the same instruments and exported to the relevant software packages
- 426 indicated below.
- 427 pKa determination was carried out by UV-Vis spectroscopy titration on $1 \times 10-3$ M solutions of the
- 428 cobalt complex, taken to 0.01 M HClO4, by adding small aliquots of 0.1 M NaOH. Electronic spectra
- 429 (Fig. S8[†]) were recorded by using a Helma 661.202-UV All Quartz Immersion Probe connected to a
- 430 Cary 50 instrument with optical fibres. The pKa determination was carried out using the standard
- 431 Specfit or ReactLab Equilibrium software.43,44
- 432

433 Kinetics

- 434 Solutions of the different ligands involved in the kinetic runs were prepared in the corresponding 0.4 M
- buffer solutions at I = 1.0 described above. The solutions of the metal complex were prepared at much
- 436 higher concentrations (20–30 fold) in water; thus an effective acidic pH was achieved, which prevented
- 437 its polymerisation processes. Small aliquots of this stock solution were added to achieve the final
- 438 conditions of the runs ([CoIII] = $(2-7) \times 10-4$ M, [ligand] = 0.01-0.1 M). For all the substitution
- 439 processes pseudo-first order flooding conditions were used.
- All the time-resolved experiments (by pH, and NMR or UV-Vis spectral monitoring) were conducted
- following three types of setups. (i) For non-buffered medium the desired aliquot of the stock CoIII
- 442 complex solution was added to a solution at a chosen acidity; pH was immediately registered and further
- 443 UV-Vis, NMR and pH changes were monitored. (ii) For experiments in buffered media the desired

- aliquot of the stock CoIII complex solution was added to the chosen 0.4 M buffer solution; pH was
- registered and further UV-Vis, NMR and pH changes were monitored. (iii) For experiments, requiring a
- 446 preequilibration process, the desired aliquot of the stock CoIII complex solution was added to the
- chosen 0.4 M buffer solution without reactants; pH was registered and UV-Vis monitoring was carried
- 448 out. When the spectral changes associated with the equilibration process were completed a solution of
- the desired ligand in the chosen 0.4 M buffer was added to the final desired concentration; pH was
- 450 registered and further UV-Vis, NMR and/or pH changes were monitored.

451 All data were collected as full (300–750 nm) spectra and treated with the standard Specfit or ReactLab

- 452 Kinetics software; 43,44 observed rate constants were obtained from the full changes of the spectra or
- alternatively at the wavelength where a maximum change was observed. The changes were fitted to the
- relevant $A \rightarrow B$ single exponential equation when first or pseudo-first order conditions applied; for
- 455 consecutive reactions with the same characteristics, an $A \rightarrow B \rightarrow C$ two exponential sequence was
- 456 fitted. Table S1[†] shows all the values obtained for kobs as a function of the different compounds and457 variables studied.
- 458

459 X-ray diffraction analyses

- 460 A red prism-like specimen of C13H26CoF9N4O12S3, with approximate dimensions $0.299 \times 0.127 \times$
- 461 0.107 mm, was used for the X-ray crystallographic analysis. The X-ray intensity data were collected on
- 462 a D8 Venture system equipped with a multilayer monochromator and a Mo microfocus ($\lambda = 0.71073$ Å).
- 463 The frames were integrated with the Bruker SAINT software package using a narrow-frame algorithm.
- 464 The integration of the data using a monoclinic unit cell yielded a total of 32 892 reflections to a
- 465 maximum θ angle of 26.40° (0.80 Å resolution), of which 5568 were independent (average redundancy
- 466 5.907, completeness = 99.6%, Rint = 4.65%, Rsig = 3.17%) and 5408 (97.13%) were greater than
- 467 $2\sigma(F2)$. The final cell constants a = 9.0812(4) Å, b = 22.5814(13) Å, c = 13.3340(7) Å, $\beta = 95.368(2)^\circ$,
- volume = 2722.4(2) Å3 are based upon the refinement of the XYZ-centroids of reflections above $20\sigma(I)$.
- 469 Data were corrected for absorption effects using the multi-scan method (SADABS). The calculated
 470 minimum and maximum transmission coefficients (based on crystal size) are 0.6885 and 0.7454.
- 471 The structure was solved using the Bruker SHELXTL software package, and refined using SHELXL,70
- 472 using the space group P121/n1, with Z = 4 for the formula unit, C13H26CoF9N4O12S3. The final
- anisotropic full-matrix leasts quares refinement on F2 with 382 variables converged at R1 = 7.27%, for
- 474 the observed data and wR2 = 16.31% for all data. The goodness-of-fit was 1.068. The largest peak in the
- final difference electron density synthesis was 1.812 e Å-3 and the largest hole was -1.239 e Å-3 with
- an RMS deviation of 0.116 e Å-3. On the basis of the final model, the calculated density was 1.851 g
- 477 cm–3 and F(000), 1544 e–.
- 478

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480

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591 Legends to figures

593	Figure 1 Drawing of the $[Co{(\mu-ET)cyclen}(H2O)2]3+$ cation prepared in this work (hydrogen atoms
594	have been omitted for clarity). Relevant distances (Å) and angles (°) are: $Co(1)-N(1) = 1.95$; $Co(1)-$
595	N(2) = 1.90; Co(1)-N(3) = 1.89; Co(1)-N(4) = 1.96; Co(1)-O(1) = 2.06; Co(1)-O(2) = 1.90; N(1)-O(2) =
596	Co(1)-N(4) = 169.0; N(2)-Co(1)-N(3) = 90.0; O(1)-Co(1)-O(2) = 85.4.
597	
598	Figure 2. (a) Changes in the electronic spectrum of a solution of $[Co{(\mu-ET)cyclen}(H2O)2]3+$
599	complex (5 \times 10–4 M) in buffered aqueous solution at Ph = 7.0 of inorganic phosphate 0.01 M (40 °C,
600	HEPES, $I = 1.0$ (NaClO4)). (b) Plot of the values of the dependence of kobs on [phosphate] at different
601	pHs at 40 °C (▲, pH = 6.0; ■, pH = 6.5; ●, pH = 7.0; 0.4 M MES/HEPES, I = 1.0 (NaClO4)).
602	
603	Figure 3 a) Changes with time of the electronic spectrum of a 5 \times 10–4 M solution of complex [Co{(μ -
604	ET)cyclen}(H2O)2]3+ with thymidine 0.04 M at $pH = 7.0$ (HEPES 0.4 M, 40 °C, $I = 1.0$ (NaClO4)). (b)
605	Plot of the values of the dependence of kobs on [thymidine] at different pHs at 40 °C (\blacktriangle , pH = 6.0; \blacksquare ,
606	$pH = 6.5; \bullet, pH = 7.0; 0.4 \text{ M HEPES}, I = 1.0 (NaClO4)).$
607	
608	Fig. 4 Plot of the values of the dependence of kobs on $[5'-TMP-/5'-TMP2-]$ at different pHs at 40 °C
609	(▲, pH = 6.2; ■, pH = 6.5; ◆, pH = 6.8; •, pH = 7.0; 0.4 M MES/HEPES, I = 1.0 (NaClO4)).

- 612
- 613





{(µ-ET)cyclen}

 $[\text{Co}\{(\mu\text{-ET})\text{cyclen}\}(\text{H}_2\text{O})_2]^{3^+}$







cytidine

O

thymidine

uridine







0

N

NH

O



5'-UMP

O

NH

FIGURE 1







SCHEME 3











SCHEME 5





-{(Me)₂(µ-ET)cyclen}



K_{OS} ca. 20



О́ Н

ОН



- **Table 1.** Summary of the kinetic, thermal and pressure activation parameters for the reaction of $[Co{(\mu-$
- ET)cyclen}(H2O)2]3+ with phosphate, cytidine, thymidine, 5'-CMP and 5'-TMP at different pHs (0.4
- 675 M MES/HEPES, I = 1.0 NaClO4)

Entering ligand	рн	²¹² k/M ⁻¹ s ⁻¹	ΔH [‡] /kJ mol ⁻¹	$\Delta S^{\ddagger}/J \text{ K}^{-1} \text{ mol}^{-1}$	$\Delta V^{\dagger}/cm^{3} mol^{-1}$
H2PO4-/HPO4-	6.0	$k_1 = 4.4 \times 10^{-4.4}$		Not determined	
		$k_2 = 4.8 \times 10^{-4.0}$			
	6.5	$k_1 = 4.4 \times 10^{-4.0}$			
		$k_2 = 4.0 \times 10^{-4.5}$			
	7.0	$k_1 = 5.2 \times 10^{-4a}$			
		$k_2 = 3.8 \times 10^{-4.5}$			
Cytidine	6.0	$k_1 = 12 \times 10^{-3}$		Not determined	
		$k_{-1} = 2.8 \times 10^{-6}$			
		$k_2 = 1.7 \times 10^{-3}$			
		$k_{-2} = 2.7 \times 10^{-5c}$			
	6.5	$k_1 = 1.3 \times 10^{-3}$	95 ± 1	19 ± 4	Not determined
		$k_{-1} = 4.4 \times 10^{-4.6}$	112 ± 1	50±4	
		$k_2 = 2.6 \times 10^{-3}$	100 ± 5	20 ± 16	
		$k_{o} = 3.0 \times 10^{-5}$	115 ± 9	41 ± 30	
	7.0	$k_{*} = 17 \times 10^{-3}$		Not determined	
		$k_{1} = 2.0 \times 10^{-3}$			
		$k_0 = 2.7 \times 10^{-3}$			
		$k_{-9} = 6.0 \times 10^{-5.2}$			
Thymidine	6.0, 6.5, 7.0	$k_1 = 4.5 \times 10^{-2d}$	95±4	11 ± 14	-8 ± 1"
	and and the	$k_2 = 3.2 \times 10^{-3}$	90 ± 7	-8 ± 22	4 ± 1"
5'-CMP*/5'-CMP ²⁻	6.0	$k_{\rm c} = 2.1 \times 10^{-3}$		Not determined	
		$k_{c} = 2.1 \times 10^{-4}$			
		$k_{0} = 9.9 \times 10^{-4}$			
		$k_{0} = 7.9 \times 10^{-5}$			
	6.5	$k_1 = 7.7 \times 10^{-3}$	115 + 4	78+14	Not determined
		$k_{-} = 4.8 \times 10^{-4.2}$	79 + 5	-58 + 16	
		$k_2 = 1.1 \times 10^{-3}$	71 ± 6	-78 ± 20	
		$k_{o} = 1.8 \times 10^{-4}$	129 ± 9	93 ± 30	
	7.0	$k_{\rm c} = 9.3 \times 10^{-3}$		Not determined	
		k = 9.2 × 10 ^{-4.4}			
		$k_{2} = 1.6 \times 10^{-3}$			
		$k_{0} = 2.4 \times 10^{-4.6}$			
SUTMP SUTMP	62	$k = 4.1 \times 10^{-3} f$			
		$k_{1} = 3.0 \times 10^{-4}$ g			
	6.5	$k_{1} = 4.1 \times 10^{-3} f$	43 ± 3^{h}	-160 ± 8^{h}	-15 ± 2^{4}
		$k_1 = 3.5 \times 10^{-4}$ g	105 ± 5 ^h	20 ± 14^{h}	0 ± 0.8^{4}
	6.8	$k_{1} = 4.1 \times 10^{-3} f$		Not determined	
	a human	$k_{0} = 4.2 \times 10^{-4g}$		and association of a	
	70	$k_{1} = 4.1 \times 10^{-3} f$			
	2.00	h = 47 × 10-48			

^a Limiting value, in s⁻¹; average value for all systems of $K_{CR} = 120 \text{ M}^{-1}$. ^b Concentration independent bridge formation path, see the text. ^c Reverse rate constants in s⁻¹, ^d Limiting value, in s⁻¹; $K_{CR} = 240 \text{ M}^{-1}$. ^b Determined at pH = 6.5 and 30 °C using $k_2 \times k_{\text{obsR}}(rapmdus) \sim 0.01 \text{ M}$, according to Fig. 3b. ^f Limiting pH independent value, in s⁻¹; $K_{CR} = 20 \text{ M}^{-1}$. ^f Dimiting value, in s⁻¹; average value for all systems of $K_{CR} = 35 \text{ M}^{-1}$. ^h At 0.1 M 5'TMP according to its limiting kinetics. ⁱ Determined at pH = 6.5 and 27 °C using $k \times k_{\text{obsR}}(rappendix) \sim 0.01 \text{ M}$.

Table 2 Crystal data and structure refinement for complex $[Co{(\mu-ET)-cyclen}(H2O)2](CF3SO3)3$

Empirical formula	C13H28C0F9N4O12S8
Formula weight	758.50
Tem perature	100(2) K
Wavelength	0.71073 Å
Crystal system	Monoclinic
Space group	P2 1/n
Unit cell dimensions	a = 9.0812(4); b = 22.5814(13);
	c = 13.3340(7) Å
	$\alpha = 90; \beta = 95.368(2); \gamma = 90^{\circ}$
Volume	2722.4(2) Å ³
Z	4
Density (calculated)	1.851 Mg m ⁻³
Absorption coefficient	0.985 mm ⁻¹
F(000)	1544
Crystal size	0.299 × 0.127 × 0.107 mm ³
θ range for data collection	2.368 to 26.396°
Index ranges	$-11 \le h \le 10, -28 \le k \le 28,$
	$-16 \le l \le 16$
Reflections collected	32 892
Independent reflections	5568 [R(int) = 0.0465]
Completeness to $\theta = 25.242^{\circ}$	99.7%
Absorption correction	Semi-empirical from equivalents
Max, and min, transmission	0.7454 and 0.6885
Refinement method	Pull-matrix least-squares on F ²
Data/restraints/para meters	5568/83/382
Goodness-of-fit on F ²	1.068
Final R indices $[I > 2o(I)]$	$R_1 = 0.0727, WR_2 = 0.1616$
R indices (all data)	$R_1 = 0.0747, WR_2 = 0.1631$
Extinction coefficient	n/a
Largest diff, peak and hole	1.812 and -1.239 e Å-2
OCDC code	140,0967