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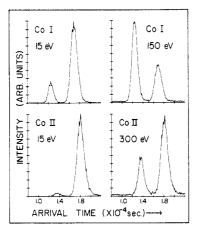


Figure 1. Arrival-time distribution of Co⁺ in He; $P_{\text{He}} \simeq 1.0$ Torr, T =160 K for CoCO₃NO (I) and CoCp(CO)₂ (II). The shift in arrival time between I and II is due to slightly different E/N values in the experiments. The times shown include time in the quadrupole, which must be subtracted to determine the mobility.

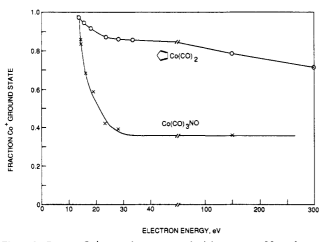


Figure 2. Percent Co+ ground state versus ionizing energy. Note change in energy axis at 50 eV.

distribution of times between injection and collection forms the arrival-time distribution. Total time between ion formation and collection is 100-400 µs. No ions other than Co⁺ and Co₋He⁺ (≲1%) were observed after the reaction cell.

Figure 1 shows the ATDs for Co+ formed from I and II. In both cases, two peaks are observed and the peak at longer time (lower mobility) increases with decreasing ionizing energy, approaching 100% at ~14 eV. Figure 2 shows the percent slower peak versus ionizing energy. An unlikely explanation is the formation of another ion with the same nominal mass which could not be differentiated in the quadrupole. The m/e 59 peak was examined at high resolution, and no such ion was found. We are convinced that the peaks are due to ground-state Co⁺(3d⁸, ³F) and a metastable excited state, probably Co⁺(3d⁷4s¹, ³F). The respective zero field reduced mobilities¹² were measured to be K_0 = 16.2 ± 2 cm²/(V s) (ground state) and K_0 = 24.1 ± 2 cm²/(V s) (excited state); their ratio is 1.485 ± 0.05 . No variation in K_0 was found between cell temperatures of 160 and 300 K.

If Co⁺ electronic states are responsible for the different mobilities, the increased mobility of the ³F state is probably due to the populated 4s orbital. The 4s orbital is considerably larger than the 3d;2,13 thus, long-range repulsion between Co+ and He should increase if it is occupied. This will reduce the effective attractive potential due to the charge-induced dipole, lowering the collision rate. Tonkyn et al.² have used this argument to help explain metal

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ion/alkane clustering data. Mobilities in helium are often determined largely by forces other than the charge-induced dipole, due to the small polarizability of helium. 12,14 The existing Ti⁺ mobility data⁹ support the above hypothesis. The Ti⁺ ground state $(3d^24s^1, {}^4F)$ includes a 4s electron⁶ and has a mobility of 25.5 \pm 0.5 cm²/(V s), 9 very similar to that of the excited-state cobalt. This may indicate that configuration (not electronic state) determines mobility. Further support for long-range repulsion between excited Co+ and He comes from the absence of deactivation in collisions with helium $(k < 1 \times 10^{-14} \text{ cm}^3/\text{s})$. Deactivation might be expected in an intimate collision. We are examining other first-row transition metals to check the correlation between high mobility and occupied 4s orbital.

Finally, we can compare the apparent effects of 3d and 4s orbital population on ion mobility. Potassium ions (3p⁶, ¹S) have a mobility of He of 21.6 cm²/(V s).^{14,15} If the metal ion/He interaction were purely electrostatic (i.e., charge-induced-dipole attraction versus electron-electron repulsion), we would expect that adding 3d electrons to K⁺ should cause some increase in mobility due to increased electron density on the ion while adding a 4s electron should cause a greater increase. 16 In fact, the 3d electrons cause a large decrease in mobility (at least for ground-state Co⁺), indicating a decreased repulsion, which must be due to other factors. Studies of other first row transition metal ions are in progress.

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Structural Studies by ¹H/¹³C Two-Dimensional and Three-Dimensional HMQC-NOE at Natural Abundance on Complex Carbohydrates

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Studies on the chain conformation of oligosaccharides are mainly restricted to analysis of NOE spectra to estimate the torsion angles defining the glycosidic linkages. 1,2 A major problem in the analysis is the overlap of resonances in the bulk region between 3 and 4 ppm. Recently, we demonstrated a homonuclear 3D NOE-HOHAHA experiment to resolve overlap in 2D NOE spectra of oligosaccharides.³ ¹³C spectra of oligosaccharides are much better resolved than ¹H spectra. Therefore, the inclusion or addition of a ¹³C frequency domain is a promising extension of 2D ¹H NOE spectroscopy. So far heteronuclear NMR experiments such as 2D and 3D HMQC-NOE have been limited to isotope-enriched proteins.⁴⁻⁹ Furthermore, ¹H/¹³C HMQC-

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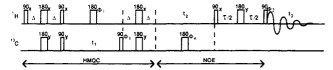


Figure 1. Pulse sequence of 3D HMQC-NOE. The following phase cycling is used: $\Phi_1 = 2(x, x, -x, -x, y, y, -y, -y); \Phi_2 = 8(x), 8(-x); \Phi_3 =$ 8(x,-x); $\Phi_4 = 4(x,-x,-x,x)$; acq = 2(x,-x), 4(-x,x), 2(x,-x). Positive and negative frequencies in ω_1 and ω_2 were separated by independent TPPI on the 90° ¹³C pulse for ω_1 and all pulses prior to the t_2 evolution period for ω_2 . The delay Δ is equal to $1/(4J(^1H^{-13}C))$; the delay τ is the NOE mixing period. The extra delays Δ after the evolution period will refocus antiphase magnetization into in-phase magnetization before starting the NOE part. The ¹H and ¹³C 180° pulses are repeated during this delay to be able to phase the spectrum afterwards. 13C decoupling in the t_2 evolution period is accomplished by a 13 C 180° pulse in the center of t_2 .

COSY and HMQC-HOHAHA have been performed. 10-13 Here we give an example of the application of ¹H/¹³C HMQC-NOE at natural abundance to the diantennery asparagine-linked oligosaccharide 1 as a tool for structural analysis.

A 3D HMQC-NOE experiment can be regarded as a combination of a 2D HMQC and a 2D NOE experiment (Figure 1). The HMQC part, containing an effective suppression of ¹²C magnetization terms, is essentially as used before. 14-16 The excitation of ¹²C z magnetization during the NOE part can be effectively suppressed by using a $90^{\circ}_{x}-\tau/2-180^{\circ}_{y}-\tau/2-90^{\circ}_{x}$ mixing sequence. 13C decoupling in the acquisition period is not necessary. The NOE magnetization exchange involves mainly transfer from a proton attached to ¹³C to a proton attached to ¹²C; by consequence, NOE cross peaks ($\omega_2 \neq \omega_3$) are not split by the ¹H-¹³C coupling. Only cross peaks containing magnetization not transferred during the NOE period ($\omega_2 = \omega_3$) will be split by the ${}^{1}H^{-13}C$ coupling in the ω_3 domain. These HMQC cross peaks are found on the diagonal ($\omega_2 = \omega_3$) of a cross-sectional plane perpendicular to the ω_1 axis. NOEs are found on a line perpendicular to the ω_2 axis. The absence of decoupling yielded a more stable measurement.

In Figure 2, three ω_1 cross sections of the 3D HMQC-NOE spectrum of 1 are shown. The small number of 112 increments in the ${}^{13}C$ t_1 direction results in a poor resolution; therefore, both GlcNAc-5/5' H-1 (δ C-1 = 100.6 ppm) and Man-4 H-1 (δ C-1 = 100.8 ppm) are present on the diagonal (Figure 2A). Comparison with a 2D NOE spectrum (insert) demonstrates that the HMQC sequence has selected a small part of the 2D NOE spectrum. The NOE effect between Man-3 H-3 and Man-4 H-1 cannot be unambiguously identified in 2D NOE nor in the homonuclear 3D NOE-HOHAHA experiment³ because of overlap with Man-4 H-5 (see Figure 2A). In 3D HMQC-NOE (Figure 2B), this NOE is not disturbed by overlap. The ω_1 cross section at $\omega_1 = 70$ ppm of NeuAc H-7 and H-4 and Gal H-4 shows an

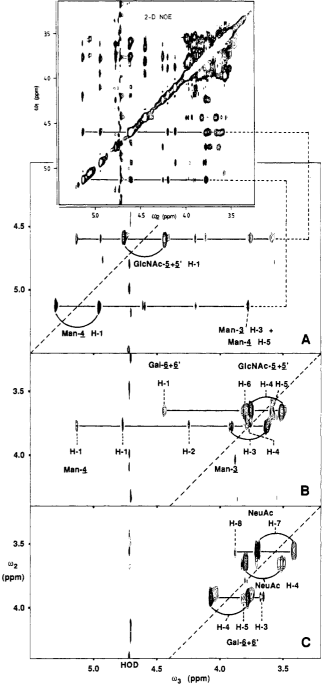


Figure 2. Cross sections perpendicular to the ω_1 axis of the 3D HMQC-NOE ¹H/¹³C spectrum of a 20 mM solution of 1 in D₂O at 304 K and pH 7. Cross sections are shown at ω_1 frequencies of 101 ppm (A), 82 ppm (B), and 70 ppm (C). The phase-sensitive spectrum was recorded at 500 MHz on a Bruker AM500 spectrometer. Acquisition was preceded by four dummy scans. FIDs of 16 scans were recorded at a size of 1K. This was repeated independently for 112 incremental t_1 and 96 incremental t_2 values, resulting in a total measuring time of approximately 63 h. The relaxation delay was 0.6 s; Δ was 1.78 ms. Spectral width in the ¹H time domains was 2272.7 Hz and, in the ¹³C time domain, 11 364 Hz. The NOE mixing time was 0.35 s. Data were zerofilled twice in the t_1 and t_2 domains before Fourier transformation. A cosine window multiplication was used in the t_1 domain, while a Hamming window was used in the t_2 and t_1 domains.¹⁷ The resulting data set of 256 × 256 × 512 points was base-line-corrected in all frequency domains by a fourth-order polynomial fit.

example of NOEs between protons resonating within the bulk region (Figure 2C). The NOE between NeuAc H-7 and H-8 is of importance to determine the conformation of the glycerol side chain of NeuAc. This NOE cannot be identified in the homonuclear 3D NOE-HOHAHA spectrum, since the small coupling

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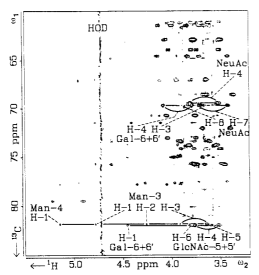


Figure 3. The bulk region of a 2D HMQC-NOE ¹H/¹³C spectrum of a 20 mM solution of 1 in D_2O at 304 K and pH 7. The phase-sensitive spectrum was recorded at 500 MHz on a Bruker AM500 spectrometer. Acquisition was preceded by four dummy scans; 2048 t_1 increments of 48 scans each were recorded at a size of 2 K, resulting in a total measuring time of approximately 60 h. The relaxation delay was 1.0 s; Δ was 1.78 ms. Spectral width in the ¹H time domains was 2272.7 Hz and, in the ¹³C time domain, 11 364 Hz. The NOE mixing time was 0.35 s. Cosine multiplications were used in the t_2 and t_1 domains. The resulting data set after Fourier transformation of 1024 × 1024 points was baseline-corrected in both frequency domains by a fourth-order polynomial

between NeuAc H-6 and H-7 prevents magnetization transfer by the isotropic mixing to the well-resolved NeuAc H-3 protons.

The extra ${}^{1}H$ dimension t_2 will separate protons attached to carbon atoms having the same ¹³C chemical shift. If there is no overlap in the ¹³C spectrum, this extra dimension is not necessary. The pulse sequence of 2D HMQC-NOE is identical with the 3D sequence except that the evolution period t_2 and the decoupling pulse have been left out. Most NOEs found in the 3D spectrum of 1 are also identified in the 2D spectrum of 1 (Figure 3). NOEs as discussed for the 3D spectrum in Figure 2, parts B and C, are indicated in Figure 3. For compounds with a well-resolved ¹³C spectrum, the 2D HMQC-NOESY sequence is a good alternative for the 3D sequence. For the majority of NOEs in the 3D spectrum of 1, the third dimension is only needed to compensate for the poor resolution in the ¹³C direction as is shown for the GlcNAc-5/5' H-4 and Man-3 H-3 in Figure 2B. With the higher resolution of the 2D HMQC-NOE (2048 t_1 increments) as compared to the 3D experiment (112 t_1 increments), the ¹³C frequencies of GlcNAc-5/5' C-4 and Man-3 C-3 are now resolved (Figure 3). The NOE between Gal-6 H-4 and Gal-6 H-5 as indicated in Figure 2C is an example of a cross peak, which cannot be identified in the 2D spectrum due to overlap of the ¹³C frequencies of Gal-6 C-4, NeuAc C-4, and NeuAc C-7. Here the extra ¹H dimension is needed to resolve the overlap by using the difference in ¹H chemical shift of Gal-6 H-4, NeuAc H-4, and NeuAc H-7 (Figure 2C).

The present work illustrates the usefulness of heteronuclear 2D and 3D NMR techniques in structure elucidations of complex oligosaccharides. Although the spectra shown are recorded at natural abundance, the sensitivity is sufficient to identify most of the NOEs of 1. Editing 2D NOE with respect to the carbon frequency gives a key to elucidating NOEs of any proton hidden in the bulk region.

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Mechanism of the C-C Cleavage of Acetone by the Ruthenium Benzyne Complex (PMe₃)₄Ru(η^2 -C₆H₄): Formation and Reactivity of an Oxametallacyclobutane Complex

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The reaction of acetone with the ruthenium benzyne complex $(PMe_3)_4Ru(\eta^2-C_6H_4)$ (1) results in cleavage of a C-C bond¹ in the ketone, leading to methane and 2, an ortho-metalated enolate complex of acetophenone (Scheme I).2 We now report the generation and spectroscopic characterization of a potential intermediate in this reaction: complex 3, a rare example of an oxametallacyclobutane. This material is formed in an unusual transformation that involves overall β -migration of a phenyl group from ruthenium to carbon in a transition-metal enolate, followed by an apparent β -alkyl elimination reaction. If, g Studies of the chemistry of complex 3 show that it undergoes a number of novel reactions, including several that involve fragmentation to α -methylstyrene.

The synthesis of oxametallacyclobutane complex 3 is shown in Scheme I. Solutions of enolate 5, the expected product from the reaction of acetone with $(PMe_3)_4Ru(\eta^2-C_6H_4)$ (1), were prepared by treatment of benzyne complex 1 with Me₃NHCl (leading to cis- and trans-phenylchlororuthenium complexes 4), followed by reaction with the potassium enolate of acetone.³ Only the O-bound isomer was detected by ¹H and ³¹P{¹H} NMR spectrometry at -20 °C, in contrast to most late-transition-metal enolates, which exist as the C-bound isomer.⁴ At room temperature over the course of an hour, complex 5 rearranges by an intramolecular migration of the phenyl group to the unsaturated enolate substituent, to form oxametallacyclobutane complex 3.5,6

Complex 3 exhibits a temperature-dependent ¹H NMR spectrum, presumably due to rapid, reversible dissociation of phosphine. Decomposition of 3, due to irreversible dissociation of phosphine

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