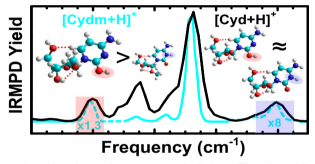


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RESEARCH ARTICLE

Structural and Energetic Effects of O2'-Ribose Methylation of Protonated Pyrimidine Nucleosides

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Abstract. The 2'-substituents distinguish DNA from RNA nucleosides. 2'-O-methylation occurs naturally in RNA and plays important roles in biological processes. Such 2'-modifications may alter the hydrogen-bonding interactions of the nucleoside and thus may affect the conformations of the nucleoside in an RNA chain. Structures of the protonated 2'-O-methylated pyrimidine nucleosides were examined by infrared multiple photon dissociation (IRMPD) action spectroscopy,

assisted by electronic structure calculations. The glycosidic bond stabilities of the protonated 2'-O-methylated pyrimidine nucleosides, [Nuom+H]⁺, were also examined and compared to their DNA and RNA nucleoside analogues via energy-resolved collision-induced dissociation (ER-CID). The preferred sites of protonation of the 2'-O-methylated pyrimidine nucleosides parallel their canonical DNA and RNA nucleoside analogues, [dNuo+H]⁺ and [Nuo+H]⁺, yet their nucleobase orientation and sugar puckering differ. The glycosidic bond stabilities of the protonated pyrimidine nucleosides follow the order: [dNuo+H]⁺ < [Nuo+H]⁺ < [Nuom+H]⁺. The slightly altered structures help explain the stabilization induced by 2'-O-methylation of the pyrimidine nucleosides.

Keywords: Cytidine (Cyd), Cytosine (Cyt), Density functional theory (DFT), Electronic structure calculations, Electrospray ionization (ESI), Energy-resolved collision-induced dissociation (ER-CID), Fourier transform ion cyclotron resonance mass spectrometer (FT-ICR MS), Gas-phase conformation, Glycosidic bond stability, Hydrogen-bonding interactions, Hydrogen-stretching region, Infrared multiple photon dissociation (IRMPD) action spectroscopy, IR fingerprint region, IRMPD spectrum, IR spectrum, 5-Methyluridine, Nucleobase, Nucleobase orientation, Nucleoside, Nucleoside modification, 2'-O-methylation, 2'-O-methylcytidine (Cydm), 2'-O-methyl-5-methyluridine (Thdm), 2'-O-methyluridine (Urdm), Protonation, Pyrimidine nucleosides, Quadrupole ion trap mass spectrometer (QIT MS), Simulated annealing, Sugar puckering, Survival yield analysis, Tandem mass spectrometry, Thymidine (Thd), Thymine (Thy), Uracil (Ura), Uridine (Urd)

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Introduction

ass spectrometry has become increasingly popular in the characterization of ribonucleosides [1–3]. Qualitative and quantitative analyses via liquid chromatography coupled with mass spectrometry (LC-MS) have contributed significantly to RNA modification analysis [4–7]. On the generic level, modifications control the activities of the biological system [8].

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During standard LC-MS analysis, RNAs are usually digested into nucleosides to facilitate analyses and achieve more consistent results. Mass spectrometry necessarily examines ions rather than neutral molecules, and protonated nucleosides are typically generated and analyzed during LC-MS analysis [4, 5, 7].

Electrospray ionization (ESI) is a soft ionization technique that typically maintains the solution-phase structures of molecules [2]. Thus, the structures of protonated nucleosides in the gas phase are commonly assumed to be the same as found from condensed-phase determinations via NMR and X-ray crystallography. However, these assumptions have generally not been validated by detailed structural investigations in the gas phase. Protonated nucleosides may adopt different sugar conformations in the gas phase due to the lack of solvent and the presence of excess charge. When digested into nucleosides, the phosphate group is cleaved off and the 3'-hydroxy substituent is also free for potential hydrogen-bonding interactions [9]. Further, the presence of the excess proton and the absence of solvent stabilizing hydrogen-bonding interactions may alter the intramolecular hydrogen-bonding interactions and even the tautomeric form of the nucleobase. The gas-phase structures of the protonated and sodium cationized forms of the canonical and some modified DNA and RNA nucleosides have been thoroughly investigated using IRMPD action spectroscopy and electronic structure methods [10-29]. The deprotonated and protonated forms of the canonical DNA and RNA mononucleotides have also been characterized by IRMPD action spectroscopy [30-40] as well as ion mobility mass spectrometry approaches [40, 41], both in conjunction with electronic structure methods. Elucidating the gas-phase structures of the protonated 2'-O-methylated pyrimidine nucleosides, [Nuom+H]⁺, may facilitate further mass spectral study of modified nucleosides and nucleic acids and provide chemical insight into the structures and folding of oligonucleotides and nucleic acids and their unimolecular decomposition behavior in the gas phase.

Modifications at the 2' position of the ribose sugar moieties may influence the overall structure and behavior of RNA. Although a large and chemically diverse set of naturally occurring modifications of RNA nucleosides are known, methylation is involved in about two thirds of the known nucleoside modifications [42-44]. Each of the canonical RNA nucleobases can undergo modification at several sites; however, modification of the ribosyl moiety has only been found to occur naturally at the 2' position [45, 46]. 2'-O-methylation is among the naturally occurring modifications, and in particular, 2'-O-methylated pyrimidines are directly involved in gene regulation, especially in the recognition of transfer-RNA (tRNA) [47–50]. For example, 2'-O-methylcytidine (Cydm) has been found in the anticodon loop of phenylalanine tRNA in yeast, 2'-O-methyuridine (Urdm) is found in glutamine tRNA of Tetrahymena and alanine tRNA of Bombyx mori [51–54], and 2'-O-methyl-5-methyluridine (Thdm) has been found in lysine and glutamic acid tRNAs from Drosophila and mammals [51, 53, 55].

Methylation of the ribosyl moiety at the 2' position leads to changes in the electronic and hydrogen-bonding characteristics of the nucleoside [9]. Compared to DNA nucleosides, the presence of the 2'-hydroxy substituent in RNA nucleosides increases the number of favorable rotational orientations of the 2'- and 3'-hydroxy substituents and the potential for canonical and noncanonical hydrogen-bonding interactions [9]. The 2'-O-methyl substituent exhibits hydrogen-bonding characteristics that are intermediate between that of DNA and RNA, as it can act as a hydrogen-bond acceptor, but not a hydrogen-bond donor. As a result, 2'-O-methyl substituents limit the nature of the hydrogen-bonding interactions possible between the 2'- and 3'-hydroxy substituents and noncanonical hydrogen bonding in some low-energy conformers of the RNA nucleosides [10–14]. Thus, the structures of the protonated forms of 2'-O-methylated nucleosides are worthy of study in their own right, but are also compared to the canonical DNA and RNA nucleosides here to elucidate the influence of 2'-O-methylation on their structures and glycosidic bond stabilities.

Structural conformations of 2'-O-methylated pyrimidine nucleosides and nucleotides have been studied using various techniques in the condensed phases [51, 56–63]. The conclusions of these studies consistently indicate that 2'-O-methyl pyrimidine nucleosides adopt 3'-endo sugar puckering and that this modification increases the structural and thermal stability of DNA and RNA strands [51, 56, 61, 64–70]. However, protonated nucleosides may adopt different sugar conformations in the gas phase due to the lack of solvent and the excess charge.

In the present work, the protonated forms of the 2'-O-methyl pyrimidine nucleosides, [Nuom+H]⁺, were investigated, including those of 2'-O-methylcytidine, [Cydm+H]⁺, 2'-O-methyluridine, [Urdm+H]⁺, and 2'-O-methyl-5-methyluridine, [Thdm+H]⁺. These nucleoside ions are examined by infrared multiple photon dissociation (IRMPD) action spectroscopy [71–74] in the IR fingerprint and hydrogen-stretching regions [75–79]. Experiments are supported by theoretical electronic structure calculations to determine the stable low-energy conformers present and their relative stabilities. Further, study of the energy-resolved collision-induced dissociation (ER-CID) behavior of these protonated pyrimidine nucleosides and their DNA and RNA analogues elucidates their relative glycosidic bond stabilities.

Experimental and Computational Methods

Materials

The 2'-O-methylated pyrimidine nucleosides, Cydm and Urdm were purchased from Alfa Aesar (Haverhill, MA, USA), whereas Thdm was purchased from Berry & Associates (Dexter, MI, USA). The canonical DNA and RNA nucleosides, and the 2'-deoxy and 5-methylated nucleosides of uridine were purchased from Sigma-Aldrich (St. Louis, MO, USA). The

HPLC grade methanol, water, and formic acid used for IRMPD experiments were purchased from Sigma-Aldrich (Saint-Quentin-Fallavier, France). The HPLC grade methanol used for the ER-CID measurements was purchased from Fischer Scientific (Waltham, MA, USA), and the HPLC grade water was purchased from Sigma-Aldrich (St. Louis, MO, USA). Ammonium acetate used in the ER-CID experiments was purchased from Fluka (St. Louis, MO, USA).

IRMPD Action Spectroscopy Experiments

The 2'-O-methylated pyrimidine nucleosides were dissolved and diluted to 1 mM in a MeOH/H₂O (50:50 v/v) mixture with 0.1% formic acid added for the IRMPD measurements in the IR fingerprint region. The IRMPD spectra of their protonated species, [Cydm+H]⁺, [Urdm+H]⁺, and [Thdm+H]⁺, were measured using a 7-T Fourier transform ion cyclotron resonance mass spectrometer (FT-ICR MS, Apex Qe, Bruker Daltonics, Bremen, Germany) coupled to the free electron laser (FEL) at the CLIO user facility [79-81]. Ions were generated via electrospray ionization (ESI), mass selected by a quadrupole mass filter, and accumulated in a hexapole ion trap prior to being transmitted to the ICR cell for photodissociation. Detailed instrumental setups have been described previously [79– 81]. The trapped [Cydm+H]⁺, [Urdm+H]⁺, or [Thdm+H]⁺ ions were irradiated for 220-390 ms with the FEL at 0-5 dB attenuation. Several (typically 3-4) mass scans were summed for IRMPD yield calculations at each IR frequency. For experiments in the hydrogen-stretching region, Cydm, Urdm, and Thdm solutions were diluted to 250 µM in a MeOH/H₂O (50:50 v/v) mixture with 0.1% formic acid added. The IRMPD spectra of [Cydm+H]⁺, [Urdm+H]⁺, and [Thdm+H]⁺ were collected on a modified quadrupole ion trap mass spectrometer equipped with an ESI source (LCQ classic, Thermo Finnigan, San Jose, CA, USA) at the University of Lyon [82–84]. Ions were irradiated by a YAG-pumped tunable IR optical parametric oscillator/amplifier (OPO) laser system (LaserVision, Bellevue, WA, USA) in the ion trap for 300-400 ms.

IRMPD spectra of $[\text{Cydm+H}]^+$, $[\text{Urdm+H}]^+$, or $[\text{Thdm+H}]^+$ were collected between $\sim 900\text{--}1900~\text{cm}^{-1}$ in the fingerprint region and $\sim 2850\text{--}3700~\text{cm}^{-1}$ in the hydrogen-stretching region. The IRMPD yield was calculated as the ratio of the sum of the intensities of fragments versus the total ion intensity, Eq. (1),

IRMPD yield =
$$\sum_{i} I_{f_i} / (\sum_{i} I_{f_i} + I_p)$$
 (1)

where I_{f_i} and I_p are the ion intensities of the fragment and precursor ions, respectively. The IRMPD yield was plotted against the laser frequency to generate the final IRMPD spectrum. Intensities were power corrected for the frequency-dependent variations in the laser output. The experimental spectra in the hydrogen-stretching region were plotted as [1-log(IRMPD yield)] to facilitate comparisons between the measured IRMPD and predicted IR spectra.

ER-CID Experiments and Survival Yield Analyses

Energy-resolved collision-induced dissociation (ER-CID) experiments were performed for the protonated forms of the DNA, RNA, and 2'-O-methylated pyrimidine nucleosides on a quadrupole ion trap mass spectrometer (OIT MS, amaZon ETD, Bruker Daltonics, Bremen, Germany) in our laboratory at Wayne State University. The ER-CID method employed has been detailed in previous work reported by our group [17–19]. Ions were generated by ESI from solutions containing 10 µM Cydm, Urdm, or Thdm in a MeOH/H₂O (50:50 v/v) mixture with 5 mM ammonium acetate. The flow rate of the syringe pump delivering the analyte solution to the ESI source was set at 3 µL/min. The ions were guided into the ion trap, and the [Cydm+H]⁺, [Urdm+H]⁺, or [Thdm+H]⁺ ions were mass isolated, accumulated, and subjected to CID in the ion trap. In addition to providing efficient trapping and cooling of the ions, the helium buffer gas (~2 mTorr for enhanced resolution mode) also served as the collision gas for the CID experiments. The q_z value of the ion trap for the ER-CID experiments was set to 0.25, which leads to a low-mass cutoff of 27% of the precursor ion m/z. The rf excitation amplitude was increased at a step size of 0.01 V from 0 V to the rf excitation amplitude required to produce complete dissociation of the precursor ions. Experiments were performed in triplicate in order to assess reproducibility. The data were processed using DataAnalysis 4.0 (Bruker Daltonics, Bremen, Germany).

The survival yield was calculated using the intensities of the precursor and fragment ions, as the ratio of the precursor ion intensity to the total ion intensity as described by Eq. (2),

Survival yield =
$$I_p / (\sum_i I_i + I_p)$$
 (2)

where I_p and I_{f_i} are defined as in Eq. (1). The survival yield was plotted as a function of the rf excitation amplitude. The rf excitation amplitude required to produce 50% dissociation of the precursor ion (CID_{50%}) was extracted by fitting the survival yield curve using the four-parameter logistic dynamic algorithm shown in Eq. (3).

$$Survival\ yield = min + \frac{(max-min)}{1 + (rf_{EA}/CID_{50\%})^{CIDslope}} \eqno(3)$$

In this equation, max and min are the maximum (1) and minimum (0) values of the survival yield, rf_{EA} is the rf excitation amplitude applied to induce fragmentation, and CIDslope is the slope of the declining region of the survival yield curve. The $CID_{50\%}$ values can be correlated with the relative glycosidic bond stabilities when glycosidic bond cleavage is the only CID pathway. $CID_{50\%}$ values of the canonical DNA, RNA, and 2'-O-methylated pyrimidine nucleosides were compared to elucidate the effects of 2'-O-methylation on their relative glycosidic bond stabilities. Data analyses were performed using SigmaPlot 10.0 (Systat Software, Inc., San Jose, CA, USA) and custom software developed in our laboratory.

Computational Details

The chemical structures of the 2'-O-methylated pyrimidine nucleosides and several stable low-energy conformers computed for their protonated forms are shown in Figure 1. The most favorable protonation sites of each 2'-O-methylated pyrimidine nucleosides were investigated. N3 and O2 protonation were examined for Cydm, whereas O2 and O4 protonation as well as 2,4-dihydroxy tautomers were considered for Urdm and Thdm based on previous investigations of the canonical DNA and RNA nucleoside analogues [12–14]. To generate candidate structures, molecular mechanics simulated annealing procedures [85] were performed using HyperChem software [86] with the AMBER 3 force field. The initial structure for each species was built by modifying (via replacement of the 2'hydroxy hydrogen atom by a methyl group) the most stable conformers determined for the analogous protonated forms of the canonical RNA nucleosides. Each initial structure was subjected to 0.3 ps of thermal heating, elevating the temperature from 0 to 1000 K. The temperature was then held constant at 1000 K for 0.2 ps to allow sampling of conformational space.

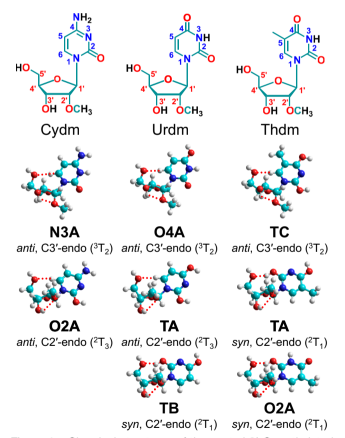


Figure 1. Chemical structures of the neutral 2'-O-methylated pyrimidine nucleosides are shown. The calculated low-energy conformers N3A and O2A of [Cydm+H]⁺, O4A, TA, and TB of [Urdm+H]⁺, and TC, TA, and O2A of [Thdm+H]⁺, optimized at the B3LYP/6-311+G(d,p) level of theory are shown. The site of protonation or tautomeric state (as indicated by the conformer designation), nucleobase orientation, and sugar puckering are given for each conformer

The system was then cooled down to 0 K over a period of 0.3 ps. The resulting structure at the end of each cycle was captured and used as the initial structure for the subsequent cycle. This process was repeated for 300 cycles for each protonation site or tautomeric form of Cydm, Urdm, and Thdm examined. Thirty low-energy structures sorted based on the molecular mechanics results were chosen for high-level quantum mechanical calculations. Additional structures were also built and examined to ensure sufficient sampling of the structure pool, and in particular, included the hydroxy and methoxy rotamers of the low-energy structures found after high level calculations. Geometry optimizations and harmonic vibrational frequency analyses were performed using the Gaussian 09 suite of programs [87]. The B3LYP/6-311+G(d,p) level of theory was employed for structural optimization and frequency analvsis to provide results directly comparable to those from previous investigations of the analogous protonated DNA and RNA nucleosides. The B3LYP and second-order Møller-Plesset perturbation with full electron correlation, MP2(full), levels of theory with a 6-311+G(2d,2p) basis set were used for single point energy calculations at 298 K, and the relative Gibbs energies of each stable conformer found were extracted and compared [10-14]. The predicted IR spectrum for each optimized structure was also extracted from the calculations. The calculated harmonic frequencies of each conformer were scaled by factors of 0.973–0.980 in the IR fingerprint region, and by 0.957-0.959 in the hydrogen-stretching region before comparison to the experimental IRMPD spectra. In order to reproduce the spectral broadening observed in the experiments, the computed IR spectra were convoluted with a Gaussian line shape having a 20 cm⁻¹ full-width-at-half-maximum (fwhm) in the IR fingerprint region and a 10 cm⁻¹ fwhm in the hydrogenstretching region.

Results

IRMPD Action Spectroscopy

The major photodissociation pathway observed for all of the protonated 2'-O-methylated pyrimidine nucleosides involves glycosidic bond cleavage with the excess proton retained by the nucleobase, resulting in elimination of the neutral 2'-O-methylated sugar moiety as summarized in reaction (4),

$$[\text{Nuom+H}]^+ + n \ hv \rightarrow [\text{Nua+H}]^+ + (\text{Nuom-Nua}) \tag{4}$$

where Nuom is Cydm, Urdm or Thdm, and Nua is the corresponding nucleobase, i.e., cytosine (Cyt), uracil (Ura), or thymine (Thy). The results for [Cydm+H]⁺ exactly parallel those found for [dCyd+H]⁺ and [Cyd+H]⁺ [12] in that reaction (4) is the only dissociation pathway observed. The results for [Urdm+H]⁺ and [Thdm+H]⁺ are comparable to those found in the IRMPD experiments of their corresponding protonated DNA and RNA nucleoside analogues, [dUrd+H]⁺, [Urd+H]⁺, [dThd+H]⁺, and [Thd+H]⁺ [13, 14]. Minor pathways involving water loss are observed in the photodissociation of [dUrd+H]⁺

and [dThd+H]⁺. Instead of water loss, the loss of methanol (M) is observed as a very minor fragmentation pathway for both [Urdm+H]⁺ and [Thdm+H]⁺ as summarized in reaction (5),

$$[\text{Nuom}+\text{H}]^+ + n \ hv \rightarrow [\text{Nuom}-\text{M}+\text{H}]^+ + \text{M}$$
 (5)

where Nuom is Urdm or Thdm. In addition, glycosidic bond cleavage resulting in loss of the neutral nucleobase from [Urdm+H]⁺ is also observed as a minor dissociation pathway as described by reaction (6), but was not observed for [dUrd+H]⁺ or [Urd+H]⁺.

$$[\text{Urdm+H}]^+ + n \text{ } hv \rightarrow [\text{Urdm-Ura+H}]^+ + \text{Ura}$$
 (6)

Due to low signal-to-noise in the experiments for [Urdm+H]⁺, the very minor dissociation pathway involving loss of methanol was not included in the IRMPD yield calculated in the IR fingerprint region, and only the dominant glycosidic bond cleavage pathway resulting in neutral loss of the sugar was included in the computed IRMPD yield in the hydrogen-stretching region. In both regions, it was confirmed that the IRMPD spectra of these minor products matched that measured for the dominant glycosidic bond cleavage pathway, and that their inclusion in the computed IRMPD yield did not alter the appearance of the spectrum except to make it noisier.

The IRMPD spectrum measured here for [Cydm+H]⁺ is compared with those measured for the corresponding canonical DNA and RNA analogues, [dCyd+H]⁺ and [Cyd+H]⁺, in Figure 2. As can be seen in the figure, the IRMPD spectra of these protonated cytosine nucleoside analogues exhibit highly parallel spectral features. One of the most obvious differences in the IRMPD spectrum of [Cydm+H]+ versus the canonical analogues is that the band at $\sim 1790 \text{ cm}^{-1}$ almost disappears. The peak at $\sim 1650 \, \mathrm{cm}^{-1}$ is also much more intense than other bands nearby; in contrast, the corresponding IR bands of [dCyd+H]⁺ and [Cyd+H]⁺ have comparable relative intensities. No IR feature is observed below 1000 cm⁻¹ for [Cydm+H]⁺, whereas two minor features are observed in the IRMPD spectra of [dCyd+H]⁺ and [Cyd+H]⁺ [12]. Splitting of the feature at ~3440 cm⁻¹ in the spectrum of [Cydm+H]⁺ parallels that observed for [Cyd+H]⁺. The two IR bands near 3550 cm⁻¹ are also fully resolved in [Cydm+H]⁺, whereas they are not resolved for the canonical DNA and RNA nucleoside analogues.

The IRMPD spectrum measured here for $[Urdm+H]^+$ is compared with those measured for the corresponding modified and canonical RNA analogues, $[dUrd+H]^+$ and $[Urd+H]^+$, in Figure 3. Only modest shifts in the band positions are observed in both the IR fingerprint and hydrogen-stretching regions. The band at $\sim 1780 \text{ cm}^{-1}$ in the spectrum of $[Urdm+H]^+$ is red shifted by 30 cm^{-1} compared to those of $[dUrd+H]^+$ and $[Urd+H]^+$. The IR feature at $\sim 1100 \text{ cm}^{-1}$ is more intense in the spectrum of $[Urdm+H]^+$ than in the spectra of $[dUrd+H]^+$ or $[Urd+H]^+$. The peak observed at $\sim 3400 \text{ cm}^{-1}$ for $[Urdm+H]^+$ is more intense than those observed for $[Urd+H]^+$ and $[dUrd+H]^+$.

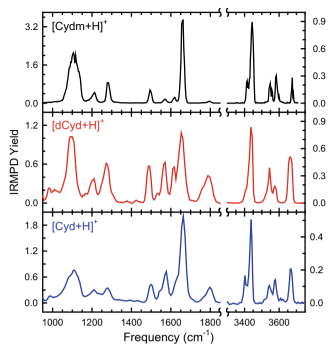


Figure 2. Comparison of the measured IRMPD spectrum of [Cydm+H]⁺ to those of its canonical DNA and RNA nucleoside analogues over the IR fingerprint and hydrogen-stretching regions. IRMPD spectra previously reported for [dCyd+H]⁺ and [Cyd+H]⁺ are taken from reference [12]

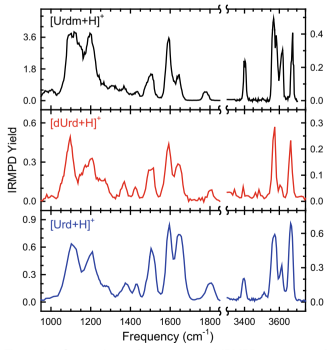


Figure 3. Comparison of the measured IRMPD spectrum of [Urdm+H]⁺ to those of its 2'-deoxy and canonical RNA nucleoside analogues over the IR fingerprint and hydrogen-stretching regions. IRMPD spectra previously reported for [dUrd+H]⁺ and [Urd+H]⁺ are taken from reference [13]

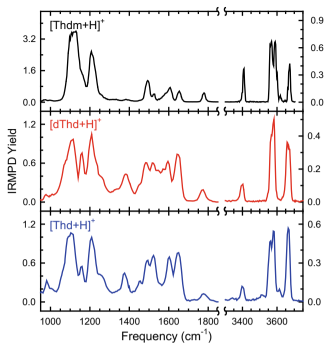


Figure 4. Comparison of the measured IRMPD spectrum of [Thdm+H]⁺ to those of its canonical DNA and 5-methylated RNA nucleoside analogues over the IR fingerprint and hydrogen-stretching regions. IRMPD spectra previously reported for [dThd+H]⁺ and [Thd+H]⁺ are taken from reference [14]

The IRMPD spectrum measured here for [Thdm+H]⁺ is compared with those measured for the corresponding canonical DNA and modified RNA analogues, [dThd+H]+ and [Thd+H]+, in Figure 4. The IRMPD spectra of these protonated thymine nucleoside analogues are also quite parallel. The most obvious spectral difference is that the band observed at $\sim 1150 \text{ cm}^{-1}$ for $[dThd+H]^{+}$ and [Thd+H]⁺ is not as well resolved, and appears as a shoulder to the blue side of the broad and intense feature at $\sim 1100 \text{ cm}^{-1}$, in the spectrum of [Thdm+H]⁺. Further, the shoulder to the blue of the intense feature at $\sim 1200 \text{ cm}^{-1}$ and the moderately intense feature observed at ~ 1380 cm⁻¹ in the spectra of [dThd+H]⁺ and [Thd+H]⁺ are much less intense in the spectrum of [Thdm+H]⁺ such that they are hardly discernible from the baseline. Peak splitting in the spectrum of $[Thdm+H]^+$ at ~3560 cm⁻¹ is more obvious than that observed in the spectra of [dThd+H]⁺ and [Thd+H]⁺. A very weak shoulder to the blue of these features is observed at ~3600 cm⁻¹ for [Thdm+H]⁺ and [Thd+H]⁺, but not $[dThd+H]^+$.

ER-CID Experimental Results

As found in the IRMPD experiments, CID of the protonated 2'-O-methylated pyrimidine nucleosides proceeds via a single major fragmentation pathway involving glycosidic bond cleavage with the excess proton retained by the nucleobase, resulting in elimination of the neutral 2'-O-methylated sugar moiety as summarized in reaction (7).

$$[Nuom+H]^{+} \stackrel{\textit{n} He}{\rightarrow} [Nua+H]^{+} + (Nuom-Nua)$$
 (7)

Representative CID mass spectra of the protonated 2'-O-methylated pyrimidine nucleosides measured at rf excitation amplitudes that result in $\sim 50\%$ fragmentation are shown in Figure 5. The results for [Cydm+H]⁺ again parallel those found for [dCyd+H]⁺ and [Cyd+H]⁺ (Figure S1), with glycosidic bond cleavage being the only fragmentation pathway observed for these protonated cytosine nucleoside analogues. Methanol (M) loss is also observed as a minor fragmentation pathway for [Urdm+H]⁺ and [Thdm+H]⁺, reaction (8). Glycosidic bond cleavage resulting in loss of the neutral nucleobase is observed for [Urdm+H]⁺ but not for [Thdm+H]⁺, reaction (9), consistent with the relative proton affinities of the Ura and Thy nucleobases [88–90].

$$[Nuom+H]^{+} \stackrel{n \text{ He}}{\rightarrow} [Nuom-M+H]^{+} + M$$
 (8)

$$[\text{Urdm} + \text{H}]^{+} \stackrel{\text{n He}}{\rightarrow} [\text{Urdm} - \text{Ura} + \text{H}]^{+} + \text{Ura}$$
 (9)

Representative CID mass spectra for the protonated uracil and thymine nucleoside analogues are compared in Figures S2 and S3. The CID fragmentation behavior of [Urdm+H]⁺ and [Thdm+H]⁺ differ significantly from their canonical and modified DNA analogues. Glycosidic bond cleavage resulting in loss of the neutral nucleobase is not observed for [Urd+H]⁺, and many water loss pathways are observed for [dUrd+H]⁺ and [dThd+H]⁺. Notably, water (W) loss has been replaced by methanol loss in [Urdm+H]⁺ and [Thdm+H]⁺, which suggests that the water and methanol losses arise from the ribose moieties. In the case of [dUrd+H]⁺ and [dThd+H]⁺, multiple pathways including water loss, even partial ribose loss in [dUrd+H]⁺ (resulting in the [Ura+W+H]⁺ fragment ion), and the nucleobase loss all contribute to the survival yield curve. The water loss pathways are dominant over neutral ribose loss in [dUrd+H]⁺, but not in [dThd+H]⁺. The higher degree of

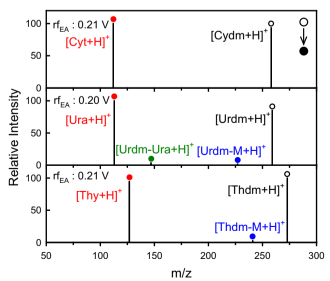


Figure 5. CID mass spectra of [Cydm+H] $^+$, [Urdm+H] $^+$, and [Thdm+H] $^+$ at rf excitation amplitudes (rf $_{EA}$) that result in $\sim 50\%$ dissociation

parallel behavior for the protonated 2'-O-methylated uracil and thymine nucleosides compared to their RNA nucleoside analogues and contrasting with that of their DNA nucleoside analogues suggest that the 2'-hydroxy and 2'-methoxy substituents stabilize the ribose moieties. Further, control experiments performed here suggest that the ketene loss pathway observed previously may have been the result of an impurity in the acetic acid used in those experiments as they were not observed in the current work [17, 18].

Survival Yield Analyses

The survival yield curves measured for the protonated 2'-O-methylated nucleosides are compared to those measured for their canonical DNA and RNA analogues in Figure 6. $CID_{50\%}$ values and estimates for the uncertainties in these values extracted from fitting the survival yield data using Eq. (3) are also shown in Figure 6. The reported uncertainties include contributions from the standard deviation in the measurements made in triplicate and the standard error of the mean associated with the least-squares fit. Trends in the $CID_{50\%}$ values of the protonated cytosine, uracil, and thymine nucleoside analogues follow the order $[dNuo+H]^+ < [Nuo+H]^+$, suggesting that 2'-O-methylation

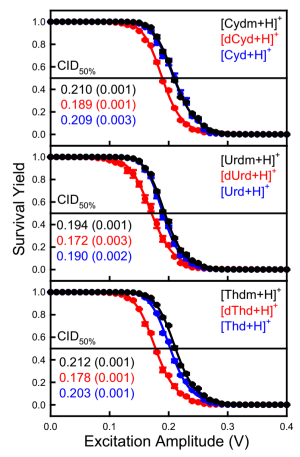


Figure 6. Comparison of the survival yield curves of $[Cydm+H]^+$, $[Urdm+H]^+$, and $[Thdm+H]^+$ to those of their canonical and modified DNA and RNA nucleoside analogues. $CID_{50\%}$ values (in V). Estimated uncertainties in these values determined from survival yield analyses are also given

strengthens the glycosidic bond relative to the protonated DNA and RNA pyrimidine nucleoside analogues. The observed trend is consistent with absolute activation energies (AEs) measured for glycosidic bond cleavage of the protonated forms of the canonical DNA and RNA pyrimidine nucleosides using threshold CID techniques [89-91]. The effect of 2'-O-methylation on the glycosidic bond stability is very slight and of the same order or smaller than the experimental uncertainties in the CID_{50%} values and thus conclusions for [Cvdm+H]⁺ are less robust. Because water losses are the dominant CID pathways for [dUrd+H]⁺ (see Figure S2), the survival yield curve and CID_{50%} value do not represent the glycosidic bond stability in this case, but instead provide a measure of the overall stability and a lower limit for the glycosidic bond stability. In contrast, the dominant CID pathways of [Urd+H]⁺ and [Urdm+H]⁺ do involve glycosidic bond cleavage. Combined with previous accurate thermochemical measurements [89], the glycosidic bond stability order: [dUrd+H]⁺ < [Urd+H]⁺ < [Urdm+H]⁺ is robust. Although water or methanol losses were observed for the protonated thymine nucleoside analogues, ribose loss still dominates the CID behavior of [dThd+H]⁺, [Thd+H]⁺, and [Thdm+H]⁺, see Figure S3. The CID_{50%} value of [Thdm+H]⁺ slightly exceeds that of [Thd+H]⁺, and is significantly higher than that of [dThd+H]+, suggesting the glycosidic bond stability follows the order $[dThd+H]^+ < [Thd+H]^+ < [Thdm+H]^+$.

The conclusions regarding the trends in the glycosidic bond stabilities based on the measured ${\rm CID}_{50\%}$ values are consistent with two-tailed t tests. The p values calculated for each pair of protonated pyrimidine nucleoside analogues are listed in Table S1. The computed p values indicate a > 95% confidence in these trends for all comparisons except $[{\rm Urd} + {\rm H}]^+$ vs $[{\rm Urdm} + {\rm H}]^+$ (> 91% confidence) and $[{\rm Cyd} + {\rm H}]^+$ vs $[{\rm Cydm} + {\rm H}]^+$ (only 60% confidence). Clearly accurate thermodynamic measurements of the absolute AEs for glycosidic bond cleavage of the protonated 2'-O-methylated pyrimidine nucleosides are of interest and would help better clarify the energetic differences that this modification produces to the stability of the protonated nucleosides.

Theoretical Results

The stable conformations of [Cydm+H]⁺, [Urdm+H]⁺, and [Thdm+H]⁺ were sampled as described in the "Computational Details" section. Each of the stable B3LYP/6-311+G(d,p) conformers optimized are classified on the basis of the site of protonation or tautomeric form, nucleobase orientation, sugar puckering, and their relative B3LYP/6-311+G(2d,2p) Gibbs energy at 298 K. The B3LYP and MP2(full) methods favor different ground-state structures and protonation sites or tautomeric forms of the protonated canonical pyrimidine nucleoside in several cases [12–14]. In the current work, relative Gibbs energies computed at the MP2(full)/6-311+G(2d,2p) level of theory are also provided for more comprehensive evaluation of the energetic trends. The classifications employed for the nucleobase orientation and sugar puckering are described in Figure S4 of the supplementary material [92, 93]. The nucleobase orientation, pseudorotation angle (P), and sugar puckering are calculated based on the methods described

previously [15–19, 63]. It is not too surprising that the structures found parallel those of the canonical pyrimidine nucleoside analogues, but exhibit slightly different sugar puckering preferences. The **N3A** conformer of [Cydm+H]⁺, Figure 1, is the lowest energy conformer found among all N3 protonated structures computed. The T conformers of [Urdm+H]⁺ or [Thdm+H]⁺ represent the 2,4-dihydroxy tautomers, and **TA** is the lowest energy conformer found among these structures. Tables 1, 2, and 3 list the relative Gibbs energies of select stable low-energy conformers calculated at the B3LYP/6-311+G(2d,2p) and MP2(full)/6-311+G(2d,2p) levels of theory for [Cydm+H]⁺, [Urdm+H]⁺, and [Thdm+H]⁺, respectively. The pseudorotation angle (*P*), nucleobase orientation, and sugar puckering of each structure are also listed.

Stable Conformations of [Cydm+H]⁺

Optimized stable B3LYP/6-311+G(d,p) N3 and O2 protonated conformers computed for [Cydm+H]⁺ are displayed in Figures 1, S5, and S6. The computed results for [Cvdm+H]⁺ are consistent with previous findings for [dCyd+H]⁺ and [Cyd+H]⁺, where B3LYP calculations favor N3 protonation over O2, and MP2 calculations suggest a preference for O2 protonation over N3 [12]. However, the energy difference predicted by MP2 for the N3A and O2A conformers is much smaller than that predicted by B3LYP (Table 1). Both N3 and O2 protonated conformers exhibit a preference for an anti nucleobase orientation, whereas conformers with a syn nucleobase orientation are predicted to be less stable. There is an obvious preference for C2'-endo or C3'-endo sugar puckering among all stable structures computed for [Cydm+H]⁺, with C3'-endo preferred over C2'-endo for the N3 protonated species by ~ 6 kJ/mol, and C2'-endo puckering slightly favored by ~ 1 kJ/mol over C3'-endo for the O2 protonated conformers.

Several conformers adopt C3'-exo or C4'-exo sugar puckerings such as conformers **N3G** and **O2I**, see Table 1 and Figures S5 and S6. It is not too surprising that these less common sugar puckerings are observed as they are the most similar to C2'-endo and C3'-endo conformations, and characterized by only a modest change in the pseudorotation angle, see Figure S4.

Stable Conformations of [Urdm+H]⁺

Optimized stable B3LYP/6-311+G(d,p) O2 and O4 protonated conformers as well as 2,4-dihydroxy tautomers (T) computed for [Urdm+H]⁺ are shown in Figures 1 and S7–S9. The low-energy structures found are quite parallel to those previously found for [dUrd+H]⁺ and [Urd+H]⁺ [13]. B3LYP exhibits a preference for O4 protonation, whereas MP2 predicts 2,4-dihydroxy tautomers as being more stable (Table 2). Interestingly, O2 protonated conformers of [Urdm+H]⁺ are just slightly less energetically favorable, by 8.3 kJ/mol for B3LYP and 5.8 kJ/mol for MP2, compared to the most stable conformers computed. The differences in stability computed for the lowest energy O2 protonated conformers of [dUrd+H]⁺ and [Urd+H]⁺ and the corresponding ground conformers are similar or somewhat larger and exceed 7 kJ/mol. The O4 protonated conformers exhibit a preference for an anti nucleobase orientation, whereas the O2 protonated conformers are stabilized by a strong O2H⁺···O5' hydrogen-bonding interaction such that the syn orientation is preferred over anti for these conformers. B3LYP predicts that the 2,4-dihydroxy tautomers prefer an anti orientation, whereas MP2 predicts that the 2,4dihydroxy tautomers prefer a syn orientation. Several O2 protonated conformers and 2,4-dihydroxy tautomers exhibit a strong O2H···O2' hydrogen-bonding interaction, and these conformers are important in the results for [Urd+H]⁺. Such O2H···O2' hydrogen-bonding interactions in [Urdm+H]⁺ disrupt or at least

Table 1. Relative Gibbs Energies of the Select Stable Low-Energy Conformers of [Cydm+H]⁺ at 298 K in kJ/mol. The Pseudorotation angles (*P*) in Degrees, Pyrimidine Nucleobase Orientations, and Sugar Puckering are also Listed

Conformer	ΔG_{298} (B3LYP)	ΔG_{298} MP2(full)	$P(^{\circ})$	Base orientation	Sugar puckering
N3A	0.0	0.5	13.2	anti	C3'-endo (³ T ₂)
N3B	6.0	6.0	168.6	anti	C2'-endo $(^2T_3)$
O2A	6.7	0.0	167.9	anti	C2'-endo $(^2T_3)$
O2B	7.9	1.4	13.6	anti	C3'-endo $(^3T_2)$
O2C	9.9	2.3	155.2	syn	C2'-endo $(^2T_1)$
N3C	10.9	11.7	169.7	anti	C2'-endo $(^2T_3)$
O2D	13.4	6.3	171.2	anti	C2'-endo $(^2T_3)$
N3D	15.3	15.5	167.7	anti	C2'-endo $(^2T_3)$
O2E	15.6	11.3	14.2	anti	C3'-endo $(^3T_2)$
O2F	17.9	10.4	154.7	syn	C2'-endo $(^2T_1)$
O2G	18.2	11.2	181.8	anti	C3'-exo $({}_{3}T^{2})^{1}$
O2H	19.5	13.5	12.0	anti	$C3'$ -endo ($^{3}T_{2}$)
N3F	22.6	22.7	159.2	syn	C2'-endo $\binom{2}{1}$
O2I	23.5	20.8	41.1	syn	$C4'$ -exo $(_4T^3)$
N3G	24.3	28.4	212.5	anti	$C3'$ -exo $({}_{3}T^{4})$
O2V	33.9	36.5	99.8	anti	O4'-endo (°T ₁)
N3P	35.4	40.6	44.4	syn	$C4'$ -exo $(_4T^3)$
O2AB	35.9	27.2	340.2	syn	$C2'$ -exo $({}_{2}T^{1})$
O2AC	36.8	37.3	143.6	anti	C1'-exo $(_1T^2)$
O2AI	49.6	46.5	38.1	anti	C4'-exo $(_4T^3)$
O2AK	58.6	59.5	6.0	syn	C3'-endo ($^{3}T_{2}$)

Table 2. Relative Gibbs Energies of the Select Stable Low-Energy Conformers of [Urdm+H]⁺ at 298 K in kJ/mol. The Pseudorotation Angles (*P*) in Degrees, Pyrimidine Nucleobase Orientations, and Sugar Puckering are also Listed

Conformer	$\Delta G_{298} \ B3LYP$	ΔG_{298} MP2(full)	$P(^{\circ})$	Base orientation	Sugar puckering
O4A	0.0	5.3	12.4	anti	C3'-endo (³ T ₂)
TA	3.2	0.8	168.5	anti	C2'-endo $(^2T_3)$
TB	3.3	0.0	155.8	syn	C2'-endo $(^2T_1)$
TC	4.5	0.5	171.9	anti	$C2'$ -endo $(^{2}T_{3})$
TD	5.1	1.9	174.7	anti	$C2'$ -endo ($^{2}T_{3}$)
TE	5.1	3.1	13.0	anti	C3'-endo $(^3T_2)$
O4B	7.1	12.0	169.1	anti	$C2'$ -endo ($^{2}T_{3}$)
O2A	8.3	5.8	157.3	syn	$C2'$ -endo $(^2T_1)$
O4C	11.3	17.2	10.5	anti	$C3'$ -endo ($^{3}T_{2}$)
O4D	11.7	18.0	170.2	anti	C2'-endo $(^2T_3)$
O4E	12.2	18.0	12.3	anti	C3'-endo $(^3T_2)$
O2B	12.3	9.7	174.7	anti	C2'-endo $(^2T_3)$
O2C	13.3	10.2	169.8	anti	C2'-endo $(^2T_3)$
O2D	16.2	13.8	157.1	syn	C2'-endo $(^2T_1)$
TF	17.0	19.1	40.1	syn	$C4'$ -exo $(_{4}T^{3})$
TG	18.9	17.0	168.4	anti	C2'-endo $(^{2}T_{3})$
TH	19.4	17.7	12.8	anti	C3'-endo (${}^{3}T_{2}$)
O2E	20.1	22.4	35.6	syn	C3'-endo ($^{3}T_{4}$)
O2F	20.9	18.1	186.1	anti	$C3'$ -exo $({}_{3}T^{2})$
TI	22.0	20.4	183.9	anti	$C3'$ -exo $(_3T^2)$
O4F	23.1	29.4	170.5	anti	C2'-endo (2T_3)
O4G	23.6	32.3	212.3	anti	$C3'$ -exo $({}_{3}T^{4})$
O4H	23.8	32.0	211.0	anti	$C3'$ -exo $(_{3}T^{4})$
O2G	24.3	28.0	40.6	syn	$C4'$ -exo $\binom{3}{4}$
O4I	25.8	30.8	159.1	syn	C2'-endo $\binom{2}{1}$
TL	25.9	23.5	169.3	anti	C2'-endo ($^{2}T_{3}$)
O2J	27.1	31.7	103.6	anti	O4'-endo (${}^{o}T_1$)
TP	30.6	30.4	186.7	anti	C3'-exo $(_{3}T^{2})$
O4N	31.1	40.7	143.9	anti	$C1'$ -exo $(_1T^2)$
O2N	31.2	30.4	170.6	syn	$C2'$ -endo $\binom{2}{1}$
O2Q	32.7	32.7	171.1	syn	C2'-endo ($^{2}T_{3}$)
O4P	33.3	44.0	171.1	anti	C3'-endo (${}^{3}T_{2}$)
TT	33.9	38.5	52.6	syn	C4'-exo $(_4T^3)$
O4S	35.9	45.9	54.0	syn	C4'-exo (4E)
O2R	36.7	42.2	143.0	anti	C1'-exo $\binom{4L}{1}$
TY	40.7	41.7	172.2		C1'-exo $\binom{1}{1}$ ' C2'-endo $\binom{2}{1}$
TAK	50.0	52.1	48.9	syn	C4'-exo $(_4T^3)$
TAN	51.5	47.2	338.6	syn	$C4$ -exo $\binom{41}{4}$ $C2'$ -exo $\binom{2}{1}$
	52.7	62.2	99.6	syn	
TAO	53.8	53.0	99.6 14.7	syn	O4'-endo (${}^{o}T_1$) C3'-endo (${}^{3}T_2$)
O2AB TAP			97.9	anti	
	54.2	62.2		anti	O4'-endo (${}^{o}T_1$)
O2AC	54.8	56.4	192.9	syn	C3'-exo $(_3T^2)$
O2AF	56.6	65.7	77.2	syn	O4'-endo (${}^{\circ}T_4$)
TAU	64.0	69.2	3.9	syn	C3'-endo (³ T ₂)
O2AP	72.2	77.5	62.5	anti	$C4'$ -exo $({}_{4}T^{o})$
O2AV	82.6	91.5	73.3	anti	O4'-endo (${}^{\circ}T_4$)
O2AY	86.2	93.4	53.7	syn	$C4'$ -exo $(_4T^3)$

Energetics based on B3LYP/6-311+G(2d,2p) and MP2(full)/6-311+G(2d,2p) single point energies computed for the B3LYP/6-311+G(d,p) optimized structures including zero point energy and thermal corrections to 298 K

destabilize the O3'H···O2' hydrogen-bonding interaction in the sugar moiety. A variety of sugar puckerings are found, with the most common C2'-endo and C3'-endo puckering modes again favored over others. O4 protonated conformers exhibit a preference for C3'-endo over C2'-endo sugar puckering by more than 7 kJ/mol, whereas C2'-endo is favored by less than 2 kJ/mol over C3'-endo for the 2,4-dihydroxy tautomers, and by more than 10 kJ/mol for the O2 protonated conformers.

Stable Conformations of [Thdm+H]⁺

Optimized stable B3LYP/6-311+G(d,p) O2 and O4 protonated conformers as well as 2,4-dihydroxy tautomers computed for

[Thdm+H]⁺ are shown in Figures 1 and S10–S12. As found for the other protonated 2'-O-methylated nucleosides, the stable conformers calculated for [Thdm+H]⁺ are highly parallel to those found for [dThd+H]⁺ and [Thd+H]⁺ [14]. Although [Thdm+H]⁺ only differs from [Urdm+H]⁺ by the 5-methyl substituent, the calculated protonation preference is altered from that of [Urdm+H]⁺. Both B3LYP and MP2 predict that 2,4-dihydroxy tautomers are the most stable among the structures computed (Table 3). B3LYP suggests that the most stable O2 and O4 protonated conformers are similar in energy and within 2 kJ/mol of the ground 2,4-dihydroxy tautomer. In contrast, MP2 finds larger energy differences among these conformers, especially for the **O4A** conformer, which is predicted to be 10.0 kJ/mol less

Table 3. Relative Gibbs Energies of the Selected Stable Low-Energy Conformers of [Thdm+H]⁺ at 298 K in kJ/mol. The Pseudorotation Angles (*P*) in Degrees, Pyrimidine Nucleobase Orientations, and Sugar Puckering are also Listed

Conformer	$\Delta G_{298}~B3LYP$	ΔG_{298} MP2(full)	$P(^{\circ})$	Base orientation	Sugar puckering
TA	0.0	0.0	154.9	syn	C2'-endo (² T ₁)
O4A	1.7	10.0	13.2	anti	C3'-endo $(^3T_2)$
O2A	1.8	4.2	156.4	syn	C2'-endo $(^2T_1)$
TB	2.4	1.5	169.4	anti	$C2'$ -endo $(^2T_3)$
TC	3.3	3.2	13.5	anti	C3'-endo $(^3T_2)$
O2B	8.2	10.2	172.5	anti	$C2'$ -endo $(^2T_3)$
O2C	9.2	10.6	168.2	anti	$C2'$ -endo (2T_3)
O4B	9.2	17.5	167.8	anti	$C2'$ -endo $(^2T_3)$
TD	10.6	12.8	15.0	anti	$C3'$ -endo $(^3T_2)$
O2D	12.6	19.1	34.7	syn	C3'-endo $(^3T_4)$
TE	12.9	18.0	40.9	syn	$C4'$ -exo $(_4T^3)$
O4C	13.7	22.9	168.9	anti	C2'-endo $(^2T_3)$
O4D	14.1	22.6	13.1	anti	C3'-endo $(^3T_2)$
O2E	16.6	18.5	185.1	anti	C3'-exo $({}_{3}T^{2})$
TF	18.1	18.8	167.3	anti	$C2'$ -endo $(^2T_3)$
O2F	18.3	26.8	41.6	syn	C4'-exo $(_4T^3)$
O2G	19.0	26.7	153.4	anti	$C2'$ -endo $\binom{2}{1}$
TG	19.0	19.0	13.4	anti	$C3'$ -endo $(^3T_2)$
O2H	19.9	25.0	170.2	syn	C2'-endo $(^2T_3)$
TI	21.6	23.5	186.4	anti	C3'-exo $({}_{3}T^{2})$
TL	23.7	25.1	165.6	anti	C2'-endo $(^2T_3)$
O4F	25.7	34.4	159.8	syn	C2'-endo $(^2T_1)$
O2N	26.8	31.5	170.9	syn	C2'-endo $(^2T_3)$
TAL	38.8	39.5	14.3	anti	C3'-endo $(^3T_2)$
O4I	39.5	53.9	43.6	syn	C4'-exo $(_4T^3)$
O2U	40.4	53.1	100.3	anti	O4'-endo (°T ₁)
TBG	47.5	57.6	99.5	anti	O4'-endo (°T ₁)
O2W	48.2	51.1	15.8	anti	C3'-endo $(^3T_2)$
TBR	54.1	60.5	128.0	syn	$C1'$ -exo $(_1T^2)$
O4P	59.5	77.1	135.7	anti	$C1'$ -exo $(_1T^2)$
O4T	71.5	82.8	149.0	syn	$C2'$ -endo $(^2T_1)$
O2AR	79.5	92.8	74.1	syn	O4'-endo (°T ₄)

Energetics based on B3LYP/6-311+G(2d,2p) and MP2(full)/6-311+G(2d,2p) single point energies computed for the B3LYP/6-311+G(d,p) optimized structures including zero point energy and thermal corrections to 298 K

stable than the TA conformer. Parallel to the results found for [Urdm+H]⁺, several O2 protonated conformers and 2,4-dihydroxy tautomers exhibit a strong O2H···O2' hydrogen-bonding interaction. In contrast to that found for [Thd+H]⁺, conformers exhibiting a O2H···O2' hydrogen-bonding interaction are less energetically favored for [Urdm+H]⁺, especially for the O2 protonated conformers, which is probably related to the disruption of the O3'H···O2' hydrogen-bonding interaction in the sugar moiety. The O2 protonated conformers and 2,4-dihydroxy tautomers exhibit a preference for a syn nucleobase orientation, whereas the O4 protonated conformers prefer an anti orientation, probably due to the lack of a strong O2H···O5' hydrogen-bonding interaction. The most common C2'-endo and C3'-endo puckering modes are again favored over all others, with C2'-endo favored over C3'endo for the O2 protonated conformers and 2,4-dihydroxy tautomers, and C3'-endo preferred over C2'-endo for the O4 protonated species.

Discussion

Conformers of [Cydm+H]⁺ Populated by ESI

The computed IR spectra for the stable conformers of $[Cydm+H]^+$ are compared with the experimentally measured

IRMPD spectrum to elucidate the structures populated in the experiments. The N3A and O2A conformers are found to match the experimental spectrum very well, see the comparison of these spectra shown in Figure 7. The most intense feature at $\sim 1660 \text{ cm}^{-1}$ in the IR fingerprint region is associated with C-N stretching and is not diagnostic as it is predicted for both O2 and N3 protonated conformers. N3 protonated conformers provide the signature carbonyl stretch at ~ 1800 cm⁻¹ as well as the major feature at $\sim 1660 \text{ cm}^{-1}$ and two small features at $\sim 1610 \text{ cm}^{-1}$ and 1560 cm⁻¹, which are related to the C–C or C-N stretches of the cytosine nucleobase. N3A also reproduces the band at $\sim 1280 \text{ cm}^{-1}$ very well. In the hydrogen-stretching region, the shoulder to the red of the strong band at ~3420 cm arises from the N3-H stretch of N3 protonated conformers. **O2A** provides the IR feature at ~ 1490 cm⁻¹, which all N3 conformers fail to reproduce. The broad feature observed at $\sim 1100 \text{ cm}^{-1}$ is associated with sugar ring breathing, and the shape of the band in the predicted IR spectrum of O2A reproduces this feature better than the N3A conformer. The weak feature predicted for N3A next to the moderately intense feature at 3585 cm⁻¹ represents the O3'-H stretch in the strong O3'H···O2' hydrogen-bonding interaction. The vibrational frequencies of such hydrogen-bonded modes are often overestimated in harmonic calculations [94–97]. The strong vibrational band 3585 cm⁻¹ is related to the O2-H

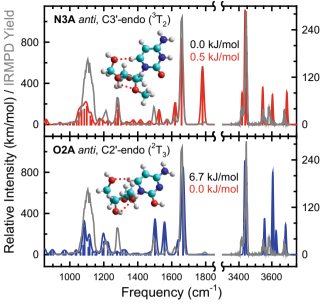


Figure 7. Comparison of the experimental IRMPD spectrum of [Cydm+H]⁺ with IR spectra predicted at the B3LYP/6-311+G(d,p) level of theory for the **N3A** and **O2A** conformers. The structures, nucleobase orientations, sugar puckering, and B3LYP/6-311+G(2d,2p) (black) and MP2(full)/6-311+G(2d,2p) (red) relative Gibbs energies at 298 K are also listed

stretch of the nucleobase calculated for O2 protonated conformers, which is slightly blue shifted compared to the measured peak. Figures S13 and S14 show similar comparisons for the lowenergy N3 and O2 protonated conformers with different sugar puckerings and nucleobase orientations. All of these latter structures exhibit spectral misalignments compared to the experimental spectra such that none of them appear to be significantly populated in the experiments. Detailed vibrational mode assignments based on the N3A and O2A conformers are listed in Table 4. In summary, N3 and O2 protonated conformers of [Cydm+H]⁺ coexist in the gas phase, with evidence for the N3A and O2A conformers being observed in the experiments.

Table 4. Vibrational Band Assignments for [Cydm+H]

Vibrational frequency (cm ⁻¹)	N3A	O2A
1102	Sugar ring stretching	Sugar ring stretching
1213	_	Sugar ring C-H bending
1277	Sugar ring C–H bending	_
1494	C4–N3 stretching	C2–N3/C4–N4 stretching
1562	C4-C5/N1-C6 stretching	C2–O stretching
1612	NH ₂ scissoring	NH ₂ scissoring
1658	C-N stretching	C-N stretching
1787	C2 = O2 stretching	_
3417	N3–H stretching	_
3445	N4H ₂ symmetric stretching	N4H ₂ symmetric stretching
3548	N4H ₂ asymmetric stretching	_
3561	_	N4H ₂ asymmetric stretching
3585	O3'-H stretching	O2-H/O3'-H stretching
3676	O5'-H stretching	O5'-H stretching

Vibrational band assignments are based on the N3A and O2A conformers

Conformers of [Urdm+H]⁺ Populated by ESI

The experimental spectrum of [Urdm+H]⁺ is compared with the IR spectra predicted for the O4A, TA, and TB conformers in Figure 8. Figure S15 shows similar comparisons for other conformers (O4F, O4G, O4H, TL, TP, O2N, O2O, O4P, TY, and TAK) that may be populated in the experiments. However, these latter conformers are less energetically favorable, and do not provide any unique features that can definitively demonstrate their presence in the experiments. The 2.4-dihydroxy tautomers generally reproduce the IR bands at ~1640 cm⁻ $\sim 1500 \text{ cm}^{-1}$, and in the region of $\sim 1050-1250 \text{ cm}^{-1}$ very well. In the hydrogen-stretching region, 2,4-dihydroxy tautomers also provide the IR features observed at ~3570 cm⁻¹. The TB conformer matches this feature even better than the TA conformer, even though the peak calculated at $\sim 1580 \text{ cm}^{-1}$ is slightly red shifted versus the measured feature. Both O4 and O2 protonated conformers may contribute to the carbonyl stretch observed at ~ 1780 cm⁻¹. The IR features in the IR fingerprint region are not sensitive to the difference between these two protonation sites. The band at $\sim 3400 \text{ cm}^{-1}$ may arise

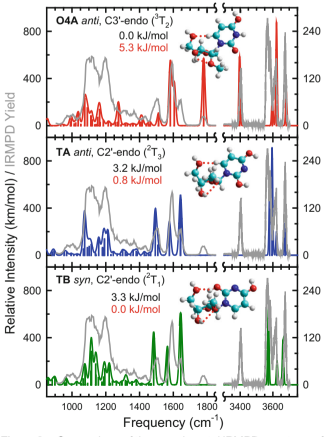


Figure 8. Comparison of the experimental IRMPD spectrum of [Urdm+H]⁺ with IR spectra predicted at the B3LYP/6-311+G(d,p) level of theory for the **O4A**, **TA**, **TB** conformers. The structures, nucleobase orientations, sugar puckering, and B3LYP/6-311+G(2d,2p) (black) and MP2(full)/6-311+G(2d,2p) (red) relative Gibbs energies at 298 K are also listed

from either O4 or O2 protonated conformers because that feature represents the N3-H stretch, which is lacking in the tautomers. However, **O2A** (Figure S18) exhibits misalignment of the IR band observed at ~3680 in the hydrogen-stretching region. The peak whose frequency is under-predicted is associated with the O5'-H stretch, which is likely the result of the hydrogen-bonding interaction with the 2-hydroxy substituent. B3LYP and MP2 both predict **O2A** as being higher in Gibbs energy compared to the **O4A** and **TA** conformers. Overall, O4 protonated conformers provide a better match to the experimental data than O2 protonated conformers, suggesting O4 protonation is more important in the experiments. Figures \$16-\$18 show comparisons of the computed IR and experimental IRMPD spectra for conformers of [Urdm+H]⁺ that adopt other nucleobase orientations and sugar puckerings. The spectral mismatches are shaded in red. It is worth pointing out that the TC and TE conformers (Figure S17) exhibit O2-H···O2' hydrogen-bonding interactions. The analogous tautomers are populated significantly in the canonical RNA nucleoside, [Urd+H]+ [13]. However, current results indicate that this conformation exhibits an obvious band shift at ~ 1450 and ~ 1550 cm⁻¹, suggesting that this conformation is not populated for [Urdm+H]⁺. The experimental IRMPD spectrum clearly arises from contributions from multiple conformers having different protonation sites or tautomeric forms (see Figure 8 and Figure S15). We conclude that the O4A, TA, and **TB** conformers are significantly populated in the experiments. Among the higher-energy conformers, O4E is the 4hydroxy rotamer of O4A. O4C also adopts a similar conformation to that of O4A, but the lack of the O3'H···O2' hydrogen bond makes it less stable than O4A. The nucleobase orientation of **O4D** is quite parallel to that of **O4A**; however, both B3LYP and MP2 find C2'-endo sugar puckering much less stable. Detailed vibrational mode assignments based on the O4A, TA, and TB conformers are listed in Table 5.

Table 5. Vibrational Band Assignments for [Urdm+H]+

Vibrational frequency (cm ⁻¹)	O4A	TA/TB
1100	Sugar ring stretching	Sugar ring stretching
1120	_	Sugar ring C–H/ scissoring/twisting
1197		O–H scissoring
1256	Sugar C-H rocking	_
1304	_	Sugar ring C-H rocking
1367	_	Sugar ring C-H wagging
1427	Sugar ring C-H/O-H scissoring	-
1500	C4–N3 stretching	C4–O4 stretching
1593	Nucleobase C-C/C-N stretching	C4–C5/C2–N2 stretching
1641	_	C5-C6/C2-N3 stretching
1773	C2=O2 stretching	-
3404	N3–H stretching	_
3567	_	O4–H stretching
3621	O4-H/O3'-H stretching	O3'-H stretching
3676	O5'-H stretching	O5'-H stretching

Vibrational band assignments are based on the O4A, TA, and TB conformers

Conformers of [Thdm+H]⁺ Populated by ESI

The measured IRMPD and the IR spectra predicted for the TA, **O2A.** and **TC** conformers of [Thdm+H]⁺ are compared in Figure 9. All three conformers have Gibbs energies within 5 kJ/mol of one other and are plausibly populated in the experiments. The TA conformer reproduces the IR feature at $\sim 1490 \text{ cm}^{-1}$ very well. The peak shapes between 1050 and 1250 cm⁻¹ also represent the experimental spectrum nicely. The **O2A** conformer provides the distinguishable carbonyl band at ~1780 cm⁻¹ in the fingerprint region and the N3-H stretch at ~3410 cm⁻¹ in the hydrogenstretching region. The TC conformer matches the strong IR band at $\sim 3590 \text{ cm}^{-1}$, which is not reproduced by either the **TA** or **O2A** conformers. Other conformers that provide a reasonable match to the experimental spectrum are shown in Figure S19. All of these latter conformers are computed to be greater than 20 kJ/mol higher in Gibbs energy than the ground conformer, and none provide any unique IR features that could confirm their presence in the experiments. Figures S20 and S21 show comparisons of the computed IR and experimental IRMPD spectra for O2 protonated conformers

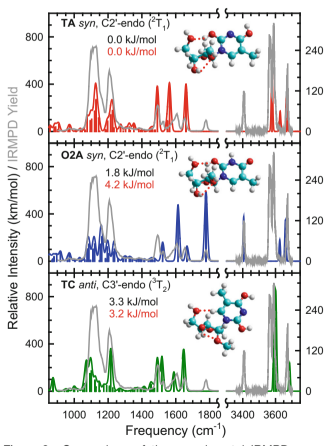


Figure 9. Comparison of the experimental IRMPD spectrum of [Thdm+H]⁺ with IR spectra predicted at the B3LYP/6-311+G(d,p) level of theory for the **TA**, **O2A**, and **TC** conformers. The structures, nucleobase orientations, sugar puckering, and B3LYP/6-311+G(2d,2p) (black) and MP2(full)/6-311+G(2d,2p) (red) relative Gibbs energies at 298 K are also listed

Table 6. Vibrational Band Assignments for [Thdm+H]⁺

Vibrational frequency (cm ⁻¹)	TA	O2A
974	C–H wagging/sugar ring bending	C–H wagging/sugar ring bending
1129	Sugar ring stretching	Sugar ring stretching
1208	C5–CH ₃ stretching/ O–H rocking	C5–CH ₃ stretching/ O–H rocking
1375	Sugar C-H twisting	_
1490	C4–O4 stretching	_
1525	_	C1-N2 stretching
1582	C2-N3/C4-C5 stretching	C2-O2 stretching
1607	C5–C6 stretching	C5–C6 stretching
1779	_	C4-O4 stretching
3407	_	N3-H stretching
3562	O4–H stretching	_
3590 ^a	_	_
3620	O3'-H stretching	O3'-H stretching
3675	O5'-H stretching	O5'-H stretching

Vibrational band assignments are primarily based on the TA and O2A conformers

and 2,4-dihydroxy tautomers that adopt other nucleobase orientations and sugar puckerings. The spectral mismatches between the predicted and experimental spectra are indicated with red shading. Consistent with [Urdm+H]⁺, conformers with an O2H···O2' hydrogen bond (TB, TBG, O2B, O2C, and O2E), cannot be significantly populated in the experiments. All O4 protonated conformers (see Figure S22) exhibit obvious shifts in the carbonyl stretch at $\sim 1780 \text{ cm}^{-1}$ and the IR band at $\sim 3590 \text{ cm}^{-1}$ that is associated with the O4–H stretch. Unlike [Urdm+H]⁺, the 5-methyl substituent makes the difference between O2 and O4 protonation more distinguishable in both regions. Overall, O2 protonated conformers and 2,4-dihydroxy tautomers are highly favored structures in the experiments. Conformers TA, O2A, and TC (see Figure 9) contribute to the experimental spectrum significantly, whereas conformers TI, TL, O2N, and TBR (Figure S19) may also contribute. Vibrational mode assignments based primarily on the TA and O2A conformers are listed in Table 6. The unique IR band contributed by the TC conformer is noted.

Influence of the O2'-Substituent on the Stable Conformations of the Protonated Pyrimidine Nucleosides

The dominant conformers N3A and O2A of [Cydm+H]⁺, and similar conformers found for [dCyd+H]⁺ and [Cyd+H]⁺ are compared in Figure S23 [12, 21]. The nucleobase always adopts an *anti* orientation, and the sugar puckering is either C2'-endo or C3'-endo. However, N3 and O2 protonated conformers prefer C2'-endo sugar puckering in [dCyd+H]⁺ and [Cyd+H]⁺. The N3A conformer of [Cydm+H]⁺ adopts C2'-endo puckering, whereas the O2A conformer adopts C3'-endo puckering. Theoretically, B3LYP always favors N3 protonation, whereas MP2 prefers O2 protonation. However, in the case of [dCyd+H]⁺ and [Cyd+H]⁺, B3LYP and MP2 results only differ by a

few kJ/mol [12]. In the case of [Cydm+H]⁺, the energy difference between N3A and O2A predicted by B3LYP is ~7 kJ/mol, whereas MP2 finds these conformers to be within 1 kJ/mol of one another, which seems more realistic on the basis of the experimental IRMPD spectrum.

According to previous work, many conformers of [dUrd+H]⁺ and [Urd+H]⁺ are populated by ESI [13]; present results suggest that even more conformers may be present in the experiments for [Urdm+H]⁺, see Figure 8 and Figure S15. Significantly populated conformers of [Urdm+H]⁺ are compared to those of [dUrd+H]⁺ and [Urd+H]⁺ in Figure S24. The O4 protonated conformers and 2,4-dihydroxy tautomers are dominant for [Urdm+H]⁺, [dUrd+H]⁺, and [Urd+H]⁺. Among all conformers populated significantly, O4 conformers always adopt an anti orientation even though the energy differences between anti and syn oriented tautomers are not large. In [dUrd+H]⁺ and [Urd+H]⁺, C2'-endo and C3'-endo sugar puckering do not cause a dramatic change in energy. For [Urdm+H]⁺, there is a clear preference for sugar puckering based on the site of protonation. The O4 protonated conformers favor C3'-endo sugar puckering (see **O4A** in Figure **S7**), and even though there is no hydrogen bond between the 2'- and 3'hydroxy substituents, the relative energy is still slightly lower than a similar conformer that adopts C2'-endo sugar puckering (compare O4C and O4D, Figure S7). This sugar puckering preference is similar to that found for the N3 protonated conformers of [Cydm+H]⁺. Interestingly, many tautomers present in the experiments adopt C2'-endo sugar puckering, whereas C3'-endo is not even found among the structures significantly populated. The sugar puckering preference is consistent with O2 protonated conformers of [Cydm+H]⁺. Lastly, the unique structures that involve O2H···O2' hydrogen-bonding interactions are only found for [Urd+H]+. Parallel conformers that have hydrogen-bonding interactions between the nucleobase 2hydroxy and sugar 2'-hydroxy substituents are found theoretically, but those structures are not populated in significant amounts in the experiments. In that regard, [Urdm+H]⁺ is more similar to [dUrd+H]⁺.

[Thdm+H]⁺, [dThd+H]⁺, and [Thd+H]⁺ all exhibit a preference for O2 protonated conformers and 2,4-dihydroxy tautomers, and their low-energy conformers are compared in Figure S25 [14, 20]. Structures with O2H···O2' hydrogen-bonding interactions are only found theoretically for [Thdm+H]+ and are predicted to be energetically competitive with other stable low-energy structures. However, comparison of the measured IRMPD and predicted IR spectra of [Thdm+H]⁺ suggest that these conformers cannot be populated significantly in the experiments, whereas they are very important for [Thd+H]⁺. Aside from these O2H···O2' hydrogen-bonded conformers, the tautomers of [Thdm+H]⁺, [dThd+H]⁺, and [Thd+H]⁺ adopt highly parallel structures. Both syn and anti orientations are found in all three species, and C2'-endo sugar puckering is favored in combination with a syn orientation. The anti oriented tautomers exhibit a preference for C2'endo sugar puckering for [dThd+H]⁺ and [Thd+H]⁺, but

 $^{^{\}rm a}\text{The IR}$ feature at $\sim\!3590~\text{cm}^{-1}$ is uniquely contributed by O4–H/O2–H stretching of the TC conformer

prefer C3'-endo sugar puckering for [Thdm+H]⁺ (see for example the **TC** conformer). For O2 protonated conformers, it is consistent that a *syn* nucleobase orientation with C2'-endo sugar puckering is favored for all three thymine nucleoside analogues, with the orientation of the hydrogen-bonding interactions in [Thd+H]⁺ and [Thdm+H]⁺ being the only difference.

Comparisons Between [Cydm+H]⁺, [Urdm+H]⁺, and [Thdm+H]⁺

The **N3A** conformer of [Cydm+H]⁺, **O4A** conformer of [Urdm+H]⁺, and **TC** conformer of [Thdm+H]⁺ all adopt *anti*, C3'-endo conformations. No *syn*, C3'-endo conformers are populated in any of these species. This is probably related to the sugar rigidity induced by the bulky 2'-O-methyl group. When the sugar adopts C2'-endo puckering, the 2'-O-methyl pyrimidine nucleosides exhibit greater flexibility in the nucleobase orientation. **O2A** of [Cydm+H]⁺ and **TA** of [Urdm+H]⁺ still adopt an *anti* orientation with ²T₃ sugar puckering. However, **TB** of [Urdm+H]⁺ and **TA** and **O2A** of [Thdm+H]⁺ adopt *syn* nucleobase orientations, and their sugar puckering changes to ²T₁. Systematic matching of the nucleobase orientation (*anti* vs. *syn*) and preferred C2'-endo puckering (²T₃ vs ²T₁) is found among all of the pyrimidine nucleosides.

Influence of the O2'-Substituent on the Dissociation Behavior of the Protonated Pyrimidine Nucleosides

As described above, the most favorable dissociation pathway for all of the protonated pyrimidine nucleoside analogues is glycosidic bond cleavage with the excess proton retained by the nucleobase, reaction (4). Based on the mechanisms and absolute energetics for glycosidic bond cleavage established for the canonical DNA and RNA nucleosides [89-91], it is clear that the proton affinities (PA) of the pyrimidine nucleobases [88] exceed those of the 2'-deoxyribose and ribose sugar moieties (although PA values have not been reported for the sugars) as glycosidic bond cleavage with the excess proton retained by the sugar, reaction (6) is more energetic by 59.8 kJ/mol for [dCyd+H]⁺, 63.7 kJ/mol for [Cyd+H]⁺, 4.8 kJ/mol for [dUrd+H]⁺, 22.1 kJ/mol for [Urd+H]⁺, 15.5 kJ/mol for [dThd+H]⁺, and 18.2 kJ/mol for [Thd+H]⁺ kJ/mol. Given the very high PA of cytosine 949.9 kJ/mol, it is not surprising that the sugar cannot compete for the excess proton and that no minor neutral loss pathways involving the sugar are observed. As the PAs of uracil and thymine are much lower, 872.7 and 880.9 kJ/mol, respectively, the sugar is able to more effectively compete for the excess proton and the neutral loss pathways become somewhat more important particularly for the dUrd and dThd. Substitution at the 2'-position stabilizes the sugar moiety such that minor neutral loss pathways involving the sugar are almost entirely shut down, but does not influence the PA of the sugar sufficiently to make glycosidic bond cleavage with the excess proton retained by the sugar, reaction (6), very competitive.

Relative Glycosidic Bond Stabilities of Protonated DNA, RNA, and 2'-O-Methylated Nucleosides

The glycosidic bond stabilities follow the order $[dNuo+H]^+ < [Nuo+H]^+ < [Nuom+H]^+$ for all of the protonated pyrimidine nucleosides, suggesting that 2'-O-methylation slightly strengthens the glycosidic bond relative to the protonated DNA and RNA nucleoside analogues. However, the guanine nucleosides were found to follow the order $[dGuo+H]^+ < [Guo+H]^+ < [Guo+H]^+$, and the glycosidic bonds of the protonated adenine nucleoside analogues are probably stronger than those of all of the other protonated nucleosides based on the significantly higher CID_{50%} values determined [24]. A detailed understanding of the energetic differences between the protonated 2'-O-methylated purine and pyrimidine nucleosides requires detailed transition state calculations and accurate guided ion beam tandem mass spectrometry measurements, which are currently being pursued by our research group.

Gas-Phase Versus Condensed-Phase Structures

All 2'-O-methylated pyrimidine nucleosides were found to prefer C3'-endo sugar puckering in the condensed phase [51, 63, 98]. The correct codon recognition and thermostability of tRNA was explained as arising from the conformational rigidity of the 2'-O-methylated nucleoside residues, in particular, the inherent stability of the C3'-endo over C2'-endo sugar puckering [51]. In the gas phase, anti, C3'-endo conformers are still present for all of the protonated 2'-O-methylated pyrimidine nucleosides, [Cydm+H]⁺, [Urdm+H]⁺, and [Thdm+H]⁺. It is not surprising that the syn or anti, C2'-endo sugar puckering conformations are competitive with the anti, C3'-endo conformations in the gas phase as a result of the flexibility and the protonation of the 2'-O-methylated pyrimidine nucleosides. On the other hand, the glycosidic bond stability of the 2'-O-methylated pyrimidine nucleosides follow the order [dNuo+H]⁺ < $[Nuo+H]^+ < [Nuom+H]^+$. The results are consistent with the increased thermostability of tRNA and other DNA and RNA studies [51, 56, 66–69].

Conclusions

IRMPD action spectroscopy experiments and theoretical analyses were performed for the 2'-O-methylated pyrimidine nucleosides to determine the structures accessed in the experiments. The preferred protonation sites or tautomeric forms of the 2'-O-methylated pyrimidine nucleosides parallel their canonical and modified DNA and RNA nucleoside analogues, yet their nucleobase orientations and sugar puckerings differ. The glycosidic bond stabilities of the protonated pyrimidine nucleosides all follow the order [dNuo+H]⁺ < [Nuo+H]⁺ < [Nuo+H]⁺ < [Nuom+H]⁺. [Cydm+H]⁺ adopts an *anti*, C3'-endo (³T₂) configuration in the most stable N3 protonated conformer, **N3A**, an *anti*, C2'-endo (²T₃) for the most stable O2 protonated conformer, **Q2A**, and these conformers dominate the experimental population. The major contributors to the measured spectrum

of [Urdm+H]⁺ are the O4 protonated, anti, C3'-endo (³T₂) conformer, O4A, and the 2.4-dihydroxy tautomers, TA and **TB** with anti, C2'-endo (${}^{2}T_{3}$) and syn, C2'-endo (${}^{2}T_{1}$) configurations, respectively. For [Thdm+H]⁺, the O2 protonated, syn, C2'-endo (${}^{2}T_{1}$) conformer, **O2A**, 2,4-dihydroxy tautomer, syn, C2'-endo (${}^{2}T_{1}$) conformer, **TA** and *anti*, C3'-endo (${}^{3}T_{2}$) conformer, TC, contribute to the IRMPD experimental spectrum. The trends in the glycosidic bond stabilities of the protonated DNA, RNA, and 2'-O-methylated pyrimidine nucleosides appear to be related to minor differences in conformation and the nature and variety of hydrogen-bonding interactions available to these species. More accurate thermochemical measurements via threshold CID approaches for the protonated forms of the 2'-O-methylated pyrimidine nucleosides are desirable and are currently being pursued to obtain absolute activation energies for glycosidic bond cleavage, and to definitively establish the influence of 2'-O-methylation on the glycosidic bond stability of the protonated pyrimidine nucleoside analogues.

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