NMR assignment of non-modified tRNA\textsuperscript{Ile} from \textit{Escherichia coli}

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Abstract
tRNAs are L-shaped RNA molecules of \textasciitilde{} 80 nucleotides that are responsible for decoding the mRNA and for the incorporation of the correct amino acid into the growing peptidyl-chain at the ribosome. They occur in all kingdoms of life and both their functions, and their structure are highly conserved. The L-shaped tertiary structure is based on a cloverleaf-like secondary structure that consists of four base paired stems connected by three to four loops. The anticodon base triplet, which is complementary to the sequence of the mRNA, resides in the anticodon loop whereas the amino acid is attached to the sequence CCA at the 3\textsuperscript{'}-terminus of the molecule. tRNAs exhibit very stable secondary and tertiary structures and contain up to 10\% modified nucleotides. However, their structure and function can also be maintained in the absence of nucleotide modifications. Here, we present the assignments of nucleobase resonances of the non-modified 77 nt tRNA\textsuperscript{Ile} from the gram-negative bacterium \textit{Escherichia coli}. We obtained assignments for all imino resonances visible in the spectra of the tRNA as well as for additional exchangeable and non-exchangeable protons and for heteronuclei of the nucleobases. Based on these assignments we could determine the chemical shift differences between modified and non-modified tRNA\textsuperscript{Ile} as a first step towards the analysis of the effect of nucleotide modifications on tRNA’s structure and dynamics.

Keywords tRNA · RNA · NMR · Assignment · Modification · Dynamics

Biological context
tRNA molecules play a central role in translation by decoding the information on the mRNA into a sequence of amino acids. During their lifetime, tRNAs encounter a variety of interaction partners: mRNAs, ribosomes, elongation factors, aminoacyl-tRNA synthetases and tRNA modification enzymes (Berg and Brandl 2021).

In all domains of life, tRNAs exhibit the same general secondary and tertiary structure. The secondary structure of tRNA is divided into five parts: the acceptor stem (Acc. stem), the dihydrouridine arm (D-arm), the anticodon-stem-loop (ACSL), the variable loop and the TΨC-arm. These elements form the well-known cloverleaf secondary structure. tRNA molecules contain a CCA-tail at the 3\textsuperscript{'}-end, where the amino acid is covalently attached to the 2\textsuperscript{'}- or 3\textsuperscript{'}-OH group. They are charged with the corresponding amino acid matching the anticodon triplet by enzymes called aminoacyl-tRNA synthetases (aaRS), yielding aminoacyl-tRNA. The recognition sites for the aaRS are often nucleotides in the acceptor stem and the anticodon loop. In three dimensional space, the tRNA molecules adopt an L-shaped tertiary structure (Kim et al. 1973). In this highly folded structure the anticodon stem and the D-arm, as well as the acceptor stem and the TΨC-arm stack coaxially. These two stacks are positioned in the L-shape through long range interactions between the D- and TΨC-loops.

tRNAs constitute the class of RNAs with the highest modification rate and more than three quarters of all known types of modified nucleotides are found within their sequences (Boccaletto et al. 2018). The extensive post-transcriptional modification modulates their overall stability and translation efficiency (Lorenz et al. 2017). However, several studies have shown that the global structure can be also achieved in unmodified tRNA with Mg\textsuperscript{2+} concentration at and above...
5 mM (Schauss et al. 2021). Therefore, it is very interesting to investigate the structural and dynamic changes that are induced by the modified nucleotides. In this respect, the structural description of modified and non-modified tRNAs is needed. Here, we present the assignments of nucleobase resonances of the non-modified 77 nt tRNA\textsuperscript{Ile} from *E. coli*.

The mature tRNA\textsuperscript{Ile} harbors total in nine modifications (Niimi et al. 1993): Dihydrouridines (D17, D20 and D20a), N6-threonylcarbamoyladenosine at position 37 (t6A37), 7-methylguanosine at position 46 (m7G46), 3-(3-amino-3-carboxypropyl)uridine at position 47 (acp3U47), ribothymidine (T54) and pseudouridines (Ψ55 and Ψ65). The comparison of chemical shifts of native and non-modified tRNA will provide the first indications of structural and dynamic changes that are induced through the modified nucleotides.

**Methods and experiments**

**Sample preparation**

The DNA template of 77 nt tRNA\textsuperscript{Ile} from *Escherichia coli* is flanked by two restriction sites and a hammerhead ribozyme:

\[\text{GAAATT (EcoRI)} - \text{TAATACGACTCACTATAG (T7 promotor)} - \text{GGGACCAAGCTTCTGATGAGTCCGTGAGG ACGAAAGACCGTCTTGGACCGTCTC (hammerhead ribozyme)} - \text{AGGCTTGTAGCTACGGTCAGGCGTACCCCTGATAAGGGTAGGTGCTGTTGTCAGTCC ACTCAGGCTTACCA (tRNA\textsuperscript{Ile})} - \text{TATG (NdeI)}.\]

Plasmid DNA and primer were purchased from Eurofins Genomics (Ebersberg, Germany). DNA was amplified by polymerase chain reaction (PCR) using T7 forward primer (5′-TAATACGACTCACTATAGG-3′) and tRNA\textsuperscript{Ile} reverse primer (5′-TGGTGCCCAGACTGCG-3′).

\[13C,15N\]-labeled 77 nt tRNA\textsuperscript{Ile} was synthesized by in vitro transcription with T7 RNA polymerase from the PCR product as described in literature (Fürtig et al. 2003; Guilleres et al. 2005). The 13C-15N-labeled (rATP, rCTP, rGTP, rUTP) nucleotides for transcription were purchased from Silantes (Munich, Germany). tRNA\textsuperscript{Ile} was purified by preparative polyacrylamide gel electrophoresis according to standard protocols (Bains et al. 2019). The tRNA\textsuperscript{Ile} was folded in water by heat denaturation for 5 min at 95 °C and immediately diluted tenfold with ice-cold water. Afterwards, tRNA\textsuperscript{Ile} was buffer-exchanged into NMR buffer (25 mM potassium phosphate, 200 mM KCl, 5 mM MgCl\textsubscript{2}, pH 6.2).

**NMR spectroscopy**

All experiments were performed in NMR buffer (25 mM potassium phosphate, 200 mM KCl, 5 mM MgCl\textsubscript{2}, pH 6.2) and 7% D\textsubscript{2}O with a sample concentration of 850 µM. All spectra were referenced to DSS (4,4-dimethyl-4-silapentane-1-sulfonic acid). Nitrogen-15 and Carbon-13 chemical shifts were indirectly referenced using the ratio of the gyromagnetic ratios of proton to 1\textsuperscript{N} (0.101329118) and 1\textsuperscript{3}C (0.251449530), respectively (Wishart et al. 1995). All experiments were performed at 25 °C if not stated otherwise.

NMR experiments were performed on 600 MHz, 700 MHz, 800 MHz, 900 MHz or 950 MHz NMR spectrometers equipped with 5 mm, z-axis gradient: 1\textsuperscript{H} \{13C, 15N\} TCI cryogenic probes (600 MHz, 800 MHz, 900 MHz, 950 MHz), a 1\textsuperscript{H} \{13C, 15N, 31P\}-QCI cryogenic probe (700 MHz) or 1\textsuperscript{3}C, 15N \{1H\}-TXO cryogenic probe (800 MHz).

NMR experiments were analyzed using Bruker Biospin software TopSpin 3.5 and assignments were performed using the software NMRFAM-SPARKY (Lee et al. 2015).

To elucidate the base-pairing interactions and assign the resonances stemming from the imino and amino groups of the nucleotides as well as other 1\textsuperscript{H}, 1\textsuperscript{5}N and 1\textsuperscript{13}C resonance from the nucleobases we conducted the following suite of NMR experiments (Table 1). To detect the homo-nuclear NOEs between imino protons and all other protons we performed a 1\textsuperscript{H}, 1\textsuperscript{H}-NOESY with jump-return water suppression (Stonehouse et al. 1994). The experiment was recorded on a 700 MHz NMR spectrometer at 25 °C with 130 ms and 170 ms mixing times and at 10 °C with 130 ms mixing time (Hwang and Shaka 1995; Sklenar 1995).

Correlation between the imino resonance and the imino nitrogen was achieved through the 1\textsuperscript{3}J_{\text{N,H}} coupling in 1\textsuperscript{H},1\textsuperscript{5}N-BEST-TROSY experiments, where the transfer delay was set to 5.4 ms. These experiments were recorded on a 700 MHz NMR spectrometer from 10 to 45 °C (Farjon et al. 2009; Favier and Brutscher 2011; Solyom et al. 2013).

Detection of correlations between fast exchanging imino and amino groups was achieved in a 1\textsuperscript{H},1\textsuperscript{5}N-CPMG-NOESY experiment, conducted with a mixing time of 130 ms (Mueller et al. 1995).

Determination of base pairing schemes in the RNA was achieved with the help of a 2D-1\textsuperscript{H},1\textsuperscript{5}N-BEST-TROSY-HNN-COSY experiment, that reports on the hydrogen bond 2\textsuperscript{3}J_{\text{N,N}}-based correlation of nitrogen atoms acting in the hydrogen bond as donor and acceptor (Dingley and Grzesiek 1998; Schulte-Herbrüggen and Sørensen 2000; Lescop et al. 2010). It was acquired with a 30 ms 1\textsuperscript{3}C,1\textsuperscript{15}N transfer delay.

Furthermore, we performed a long range 1\textsuperscript{H}, 1\textsuperscript{5}N-sfH-MQC to correlate adenine N1/N3 nitrogens with H2-protons (transfer delay 20 ms).

CN-correlations in amino groups were achieved with a 1\textsuperscript{13}C,1\textsuperscript{15}N-HSQC (Schnieders et al. 2019).

To correlate amino protons to their respective nitrogen we performed 1\textsuperscript{H},1\textsuperscript{13}C-HSQC experiments with a relaxation and exchange optimized transfer delay of 4.6 ms (Mori et al. 1995).
Correlation within aromatic CH groups was achieved with an $^1$H-$^13$C-HSQC experiment (INEPT transfer 2.7 ms) with off-resonant decoupling (Bodenhausen and Ruben 1980).

### Extent of assignment and data deposition

Before assigning the 77 nt *E. coli* tRNA$^{lle}$ (Fig. 1A), we acquired 2D-$^1$H-$^15$N-BEST-TROSY spectra at different temperatures ranging from 10 to 45 °C to elucidate at which temperature most of the exchangeable protons give rise to sharp and well dispersed signals. Furthermore, this temperature series allowed to deduce which base pairs are less stable and might stem from e.g. the closing base pairs of helices. Thereby, we found that at a temperature of 25 °C the spectra exhibited the most suitable signature of resonances to conduct assignment experiments. In total 29 well-resolved and intense imino resonances of guanosine and uridine nucleotides are visible in the spectra. All these resonances could be assigned unambiguously. Additionally, we observe four signals with a much-reduced intensity (1:10) which we attribute to a low populated second conformational state. With respect to the predicted secondary structure (Silvian et al. 1999), we could detect and assign all expected resonances of imino groups for base-paired nucleotides except for the A1:U72. The signal for this terminal base pair could be rendered beyond detectability due to dynamic motions in the ACCA-3’ end of the tRNA. Except for G19 and G46, we could detect imino proton resonances of all G and U nucleotides involved in tertiary interactions.

Table 1  List of NMR experiments with experimental parameters

|                      | Field (MHz) | Interscan delay (s) | No scans | Carrier frequencies | Spectral width | Acquisition time | Points |
|----------------------|-------------|---------------------|----------|--------------------|----------------|------------------|--------|
| $^1$H, $^1$H-NOESY   | 700         | 1.5                 | 115      | 4.7 ppm ($^1$H, $t_2$) to 7.5 ppm ($^1$H, $t_1$) | 25 ppm ($^1$H), 17 ppm ($^1$H) | 99 ms ($^1$H), 21 ms ($^1$H) | 3494 ($^1$H), 512 ($^1$H) |
| $^1$H, $^1$5N-BEST- | 700         | 0.2                 | 16       | 4.7 ppm ($^1$H), 153.5 ppm ($^{15}$N) | 25 ppm ($^1$H), 30 ppm ($^{15}$N) | 80 ms ($^1$H), 53 ms ($^{15}$N) | 2804 ($^1$H), 228 ($^{15}$N) |
| TROSY               | 600         | 1.5                 | 258      | 4.7 ppm ($^1$H), 116.0 ppm ($^{15}$N) | 25 ppm ($^1$H), 100 ppm ($^{15}$N) | 99 ms ($^1$H), 42 ms ($^{15}$N) | 2998 ($^1$H), 512 ($^{15}$N) |
| $^2$D-$^1$H, $^1$5N- | 600         | 0.3                 | 400      | 4.7 ppm ($^1$H), 185.0 ppm ($^{15}$N) | 21 ppm ($^1$H), 120 ppm ($^{15}$N) | 63 ms ($^1$H), 24 ms ($^{15}$N) | 1584 ($^1$H), 360 ($^{15}$N) |
| BEST-TROSY-HNN-COSY |             |                     |          |                    |                |                  |        |
| $^1$H, $^1$5N-NsHMQC | 800         | 1.0                 | 304      | 4.7 ppm ($^1$H), 145.0 ppm ($^{15}$N) | 10 ppm ($^1$H), 80 ppm ($^{15}$N) | 63 ms ($^1$H), 18 ms ($^{15}$N) | 1024 ($^1$H), 236 ($^{15}$N) |
| $^1$5C, $^1$5N-HSQC | 800         | 2.0                 | 512      | 4.7 ppm ($^1$H), 155.0 ppm ($^{13}$C), 85.75 ppm ($^{15}$N) | 70 ppm ($^{13}$C), 31.5 ppm ($^{15}$N) | 36 ms ($^{13}$C), 26 ms ($^{15}$N) | 1024 ($^{13}$C), 136 ($^{15}$N) |
| $^1$H, $^1$5N-HSQC  | 800         | 0.8                 | 368      | 4.7 ppm ($^1$H), 86.5 ppm ($^{13}$C), 101 ppm ($^{15}$N) | 16 ppm ($^1$H), 38 ppm ($^{15}$N) | 40 ms ($^1$H), 41.5 ms ($^{15}$N) | 1024 ($^1$H), 256 ($^{15}$N) |
| $^1$H-$^1$5C-HSQC   | 900         | 1.0                 | 512      | 4.7 ppm ($^1$H), 86.75 ppm ($^{13}$C), 143 ppm ($^{15}$N) | 8.2 ppm ($^1$H), 24 ppm ($^{13}$C) | 68 ms ($^1$H), 47 ms ($^{13}$C) | 1024 ($^1$H), 512 ($^{13}$C) |

$^1$H-$^1$5N-BEST-TROSY-HNN-COSY experiment correlated G-N1 to C-N3 and U-N3 to A-N1 and A-N7. In addition, we could assign all detectable NH$_2$ groups of paired Gs, As, and Cs through the application of an $^1$H-$^1$5N-CPMG-NOESY-HSQC (except for A67) (Fig. 1D). The protons of NH$_2$ groups were further assigned by $^1$H-$^1$5N-HSQC selective for resonances stemming from the nucleobase’s amino groups. The use of $^1$H-$^1$5N-sHMQCs allowed the correlation of adenine N1/N3 nitrogens to the H2-protons. Further, nitrogen to carbon correlations for amino groups were achieved with a $^1$3C-$^1$5N-HSQC, enabling the assignment of the C4 (for adenine and cytosine) and C2 (for guanosine) resonances. Correlation within aromatic CH groups in adenosines (C8H8 and C2H2) was achieved with a $^1$H-$^13$C HSQC experiment.

The two G-U wobble base pairs served as starting point for the resonance assignment in the 2D $^1$H-$^1$H-NOESY experiments at 10 °C and 25 °C (Fig. 1B) based on their strong intra-base pair NOEs. The imino protons of the U5:G68 base pair in the acceptor stem were assigned due to sequential NOEs to the imino proton of U6 in the U6:A67 base pair and the imino proton of G68 from the C4:G68 base pair. With this starting point the remainder of the imino protons from the other Watson–Crick base pairs in the acceptor stem were unambiguously assigned based on sequential NOEs from the other Watson–Crick base pairs in the acceptor stem. The imino protons of the G49:U65 wobble base pair at the apex of the ΨC-arm show sequential NOEs to the imino
proton of G50. With this starting point, the base paired region of the TΨC-arm was unambiguously assigned (from G49-C65 to G53-C61). Additionally, the imino groups of nucleotides U54 and U55 could be assigned. These two residues are essential for the formation of the tertiary interactions within the TΨC-loop and with residues from the D-loop. U54 forms a reverse-Hoogsteen base pair with A58 (Zagryadskaya et al. 2003) highlighted by a cross peak in the HNN-COSY spectrum between the imino proton H3 and the nitrogen N7 acting as hydrogen bond acceptor nitrogen (229.4 ppm, 15N). The hydrogen bond acceptor of U55’s imino proton is the phosphate oxygen of A58 emphasized by the lack of a cross peak in the HNN experiment and a cross peak in a long range 1H-31P-HMQC experiment (Duchardt-Ferner et al. 2011; Duchardt-Ferner and Wöhnert 2017) to a phosphate resonance. However, the peak of the imino resonance is undetectable at 45 °C in the BEST-TROSY experiment. Both, the resonances of U55 and U54 are part of the walk of NOESY connectivities in the TΨC-arm of the tRNA.

Through the stacking of the acceptor stem and the TΨC-arm (Fig. 1A), nucleotides G49 and G7 come into close proximity. This is reported by a cross peak between their two imino proton resonances. The resonance of G7 is then also the starting point for a sequential walk of NOESY cross peaks through the acceptor stem that includes base pairs G7-C66 to G2-C72. This also corroborates the before mentioned assignment of the wobble base pair G68-U5.

The imino proton resonances stemming from the D-arm were assigned starting from G22 (G22, U12, G24 and G10). The imino proton of G10 showed an additional NOE contact to G45 in the variable loop. G45 has an NOE contact to G26 that is observable as a distinct and resolved peak in the NOESY spectrum at 10 °C (Westhof and Leontis 2021).
The resonance of U8 is typically the most downfield shifted peak and was assigned at 14.27 ppm (\(^1\)H) and 164.24 ppm (\(^{15}\)N). Furthermore, its imino group is hydrogen bonded to N7 of A14, which is confirmed in the HN-COSY spectrum (Fig. 1C). The close spatial proximity of U8 to G22 and G15 induces NOE cross peaks between their imino proton resonances, exhibiting a strong NOE between U8 and G22 and a weaker NOE between U8 and G15. The remaining G-U-G-G-G NOESY walk was assigned to the anticodon-stem-loop (ACSL) consisting of the Watson–Crick base pairs G43:C27 to G39:C31 (Fig. 1A).

In summary, for the imino proton resonances, we assigned 19 out of 22 base-paired G residues (86%) and 9 out of 10 base-paired U residues (90%). The corresponding hydrogen bond acceptor nuclei C-N3, A-N1 and A-N7 were all assigned by the \(^1\)H-\(^{15}\)N-BEST-TROSY-HNN-COSY experiment. In addition, 25 amino groups were assigned (G-N2, C-N4 and A-N6, 42% in total). We assigned 28% of G-C2 and 74% of C-H41 and C-H42. Furthermore, for 4 out of 5 AU base pairs present in tRNA\(^{\text{ile}}\), we assigned the corresponding A-N3, A-C2 and A-H2 resonances.

Based on these assignments we could determine the chemical shift differences between modified tRNA\(^{\text{ile}}\) (Niimi et al. 1993) and non-modified tRNA\(^{\text{ile}}\) as a first step towards the analysis of the effect of nucleotide modifications on tRNA’s structure and dynamics. The chemical shift differences are shown in Fig. 2. The most prominent differences in chemical shifts are observed for residues close to the helix junctions (e.g. G10) and for the nucleotides that establish the tertiary interaction between the D-arm and the \(\mathrm{T}_{\Psi}\mathrm{C}\) loop, where also the nucleobase modifications T54 and \(\Psi\)55 occur. In addition, residue 65 shows a large chemical shift difference, as U65 is modified to \(\Psi\)65 in the modified tRNA\(^{\text{ile}}\). It is noteworthy to mention that the compared spectra were referenced differently, resulting in an absolute chemical shift difference of ~2.3 ppm in the \(^{15}\)N dimension. Although there are small differences in the buffer composition and for the temperature in which both tRNAs were measured, the structure of the tRNA is not perturbed and therefore chemical shifts are comparable.

The chemical shifts for tRNA\(^{\text{ile}}\) have been deposited in the Biological Magnetic Resonance Bank (http://www.bmrb.wisc.edu) with the Accession Number 51164.

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Author contributions VDJ prepared the sample. VDJ and BF performed the NMR experiments. VDJ, TB, JF and BF analyzed the data. VDJ, TB and BF wrote the manuscript. All authors provided critical comments. All authors read and approved the final manuscript.

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Data availability Chemical shift data is available at the Biological Magnetic Resonance Bank (http://www.bmrb.wisc.edu).
Declarations

Conflict of interest The authors declare that they have no conflict of interest.

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References

Bains JK, Blechar J, de Jesus V et al (2019) Combined smFRET and NMR analysis of riboswitch structural dynamics. Methods 153:22–34. https://doi.org/10.1016/j.ymeth.2018.10.004
Berg MD, Brandl CJ (2021) Transfer RNAs: diversity in form and function. RNA Biol 18:316–339
Boccaletto P, MacHnicka MA, Purta E et al (2018) MODOMICS: a database of RNA modification pathways. 2017 update. Nucleic Acids Res 46:D303–D307. https://doi.org/10.1093/nar/gkx1030
Bodenhausen G, Ruben DJ (1980) Natural abundance nitrogen-15 NMR by enhanced heteronuclear spectroscopy. Chem Phys Lett 69:185–189. https://doi.org/10.1016/0009-2614(80)80041-8
Dingley AJ, Grzesiek S (1998) Direct observation of hydrogen bonds in nucleic acid base pairs by. J Am Chem Soc 7863:714–718
Duchardt-Ferner E, Wöhnert J (2017) NMR experiments for the rapid identification of P=O···H-X type hydrogen bonds in nucleic acids. J Biomol NMR 69:101–110. https://doi.org/10.1007/S10858-017-0140-7
Duchardt-Ferner E, Ferrer J, Wöhnert J (2011) Rapid identification of noncanonical RNA structure elements by direct detection of OH⋯=O=P, NH⋯=O=P, and NH 2⋯=O=P hydrogen bonds in solution NMR spectroscopy. Angew Chem Int Ed 50:7927–7930. https://doi.org/10.1002/anie.2011101743
Farjon J, Boisbouvier J, Schanda P et al (2009) Longitudinal-relaxation-enhanced NMR experiments for the study of nucleic acids in solution. J Am Chem Soc 131:8571–8577. https://doi.org/10.1021/ja901633y
Favier A, Truttschner B (2011) Recovering lost magnetization: polarization enhancement in biomolecular NMR. J Biomol NMR 49:9–15. https://doi.org/10.1007/s10858-010-9461-5
Fürtig B, Richter C, Wöhnert J, Schwabé H (2003) NMR Spectroscopy of RNA. ChemBioChem 4:936–962. https://doi.org/10.1002/cbc.200300700
Guilleres J, Lopez PJ, Proux F et al (2005) A mutation in T7 RNA polymerase that facilitates promoter clearance. Proc Natl Acad Sci USA 102:5958–5963. https://doi.org/10.1073/pnas.0407141102
Hwang TL, Shaka AJ (1995) Water suppression that works. excitation sculpting using arbitrary wave-forms and pulsed-field gradients. J Magn Reson Ser A 112:275–279. https://doi.org/10.1007/JMRA.1995.1047
Kim SH, Quigley GJ, Suddath FL et al (1973) Three-dimensional structure of yeast phenylalanine transfer RNA: folding of the polynucleotide chain. Science 179:285–288. https://doi.org/10.1126/SCIENCE.179.4070.285
Lee W, Tonelli M, Markley JL (2015) NMRFAM-SPARKY: enhanced software for biomolecular NMR spectroscopy. Bioinformatics 31:1325–1327. https://doi.org/10.1093/BIOINFORMATICS/BTU830
Leontis NB, Westhof E (2001) Geometric nomenclature and classification of RNA base pairs. RNA 7:499–512. https://doi.org/10.1017/S1355832010002515
Lescop E, Kern T, Brutschen B (2010) Guidelines for the use of band-selective radiofrequency pulses in hetero-nuclear NMR: example of longitudinal-relaxation-enhanced BEST-type 1H–15N correlation experiments. J Magn Reson 203:190–198. https://doi.org/10.1016/j.jmre.2009.12.001
Lorenz C, Lünse CE, Mörl M (2017) Trna modifications: impact on structure and thermal adaptation. Biomolecules. https://doi.org/10.3390/biom7020035
Mori S, Abeygunawardana C, Johnson MO, Van Zijl PC M (1995) Improved sensitivity of HSQC spectra of exchanging protons at short interscan delays using a new Fast HSQC (FHSQC) detection scheme that avoids water saturation. J Magn Reson Ser B 108:94–98. https://doi.org/10.1002/jmrb.1995.1109
Mueller L, Legault P, Pardi A (1995) Improved RNA structure determination by detection of NOE contacts to exchange-broadened amino protons. J Am Chem Soc 117:11043–11048. https://doi.org/10.1021/ja00150a001
Niimi T, Takayanagi M, Naguchi T et al (1993) A 15N–1H nuclear magnetic resonance study on the interaction between isoleucine RNA and isoleucyl-1RNA synthetase from Escherichia coli. Biochimie 75:1109–1115
Schauss J, Kundu A, Fingerhut BP, Elsaesser T (2021) Magnesium contact ions stabilize the tertiary structure of transfer RNA: electrostatics mapped by two-dimensional infrared spectra and theoretical simulations. J Phys Chem B 125:740–747. https://doi.org/10.1021/acs.jpcb.0c08966
Schnieders R, Wolter AC, Richter C et al (2019) Novel 13C-detected NMR experiments for the precise detection of RNA structure. Angew Chem Int Ed 58:9140–9144. https://doi.org/10.1002/ANIE.201904057
Schulte-Herbrüggen T, Sörensen OW (2000) Clean TROSY: compensation for relaxation-induced artifacts. J Magn Reson 144:123–128. https://doi.org/10.1006/JMRE.2000.2020
Silvian LF, Wang J, Steitz TA (1999) Insights into editing from an Ile-tRNA synthetase structure with tRNAile and mupirocin. Science 285:1074–1077. https://doi.org/10.1126/SCIENCE.285.5430.1074
Sklenar V (1995) Suppression of radiation damping in multidimensional NMR experiments using magnetic field gradients. J Magn Reson Ser A 114:132–135. https://doi.org/10.1006/JMRA.1995.1119
Solyom Z, Schwarten M, Geist L et al (2013) BEST-TROSY experiments for time-efficient sequential resonance assignment of large disordered proteins. J Biomol NMR 55:311–321. https://doi.org/10.1007/s10858-013-9715-0
Stonehouse J, Shaw GL, Keeler J (1994) Improving solvent suppression in jump-return NOESY experiments. J Biomol NMR 4:799–805. https://doi.org/10.1007/BF00398410
Westhof E, Leontis NB (2021) An RNA-centric historical narrative around the Protein Data Bank. J Biol Chem 296:100555. https://doi.org/10.1016/j.jbc.2021.100555
Wishart DS, Bigam CG, Yao J et al (1995) 1H, 13C and 15N chemical shift referencing in biomolecular NMR. J Biomol NMR 6:135–140. https://doi.org/10.1007/BF00217777
Zagryadskaia EL, Doyon FR, Steinberg SV (2003) Importance of the reverse Hoogsteen base pair 54–58 for tRNA function. Nucleic Acids Res 31:3946–3953. https://doi.org/10.1093/nar/gkg448

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