Selective $^{13}$C labeling of nucleotides for large RNA NMR spectroscopy using an E. coli strain disabled in the TCA cycle

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Received: 20 May 2010 / Accepted: 28 July 2010 / Published online: 6 November 2010
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Abstract     Escherichia coli (E. coli) is an ideal organism to tailor-make labeled nucleotides for biophysical studies of RNA. Recently, we showed that adding labeled formate enhanced the isotopic enrichment at protonated carbon sites in nucleotides. In this paper, we show that growth of a mutant E. coli strain DL323 (lacking succinate and malate dehydrogenases) on $^{13}$C-2-glycerol and $^{13}$C-1,3-glycerol enables selective labeling at many useful sites for RNA NMR spectroscopy. For DL323 E. coli grown in $^{13}$C-2-glycerol without labeled formate, all the ribose carbon atoms are labeled except the C3′ and C5′ carbon positions. Consequently the C1′, C2′ and C4′ positions remain singlet. In addition, only the pyrimidine base C6 atoms are substantially labeled to ~96% whereas the C2 and C8 atoms of purine are labeled to ~5%. Supplementing the growth media with $^{13}$C-formate increases the labeling at C8 to ~88%, but not C2. Not unexpectedly, addition of exogenous formate is unnecessary for attaining the high enrichment levels of ~88% for the C2 and C8 purine positions in a $^{13}$C-1,3-glycerol based growth. Furthermore, the ribose ring is labeled in all but the C4′ position, such that the C2′ and C3′ positions suffer from multiplet splitting but the C5′ position remains singlet and the C1′ position shows a small amount of residual C1′–C2′ coupling. As expected, all the protonated base atoms, except C6, are labeled to ~90%. In addition, labeling with $^{13}$C-1,3-glycerol affords an isolated methylene ribose with high enrichment at the C5′ position (~90%) that makes it particularly attractive for NMR applications involving CH$_2$-TROSY modules without the need for decoupling the C4′ carbon. To simulate the tumbling of large RNA molecules, perdeuterated glycerol was added to a mixture of the four nucleotides, and the methylene TROSY experiment recorded at various temperatures. Even under conditions of slow tumbling, all the expected carbon correlations were observed, which indicates this approach of using nucleotides obtained from DL323 E. coli will be applicable to high molecular weight RNA systems.

Keywords     Alternate-site specific labeling · CH$_2$-methylene TROSY · Ribose and nucleobase · RNA · Structure and dynamics

Abbreviations

AMP     Adenosine 5′-monophosphate
CMP     Cytidine 5′-monophosphate
DHAP    Dihydroxyacetone phosphate
FBP     Fructose-6-bisphosphate
F6P     Fructose-6-phosphate
GA3P    Glyceraldehyde-3-phosphate
Gly     Glycine
GMP     Guanosine 5′-monophosphate
noPPP   Non-oxidative pentose phosphate pathway
OAA     Oxaloacetate
oPPP    Oxidative pentose phosphate pathway
R5P     Ribose-5-phosphate
rNMPs   Ribonucleoside monophosphates
rNTPs   Ribonucleoside triphosphates
Ser     Serine
TIM     Triosephosphate isomerase
UMP     Uridine 5′-monophosphate
Introduction

Uniform labeling technology ushered in an era of high resolution NMR studies of RNA structures and RNA-ligand interactions (Batey et al. 1995; Hall 1995; Pardi 1995; Fürtig et al. 2003; Latham et al. 2005; Dayie 2008; Lu et al. 2010). Milligram quantities of uniformly $^{13}$C/$^{15}$N labeled nucleotides can be extracted from bacteria grown on minimal media supplemented with $^{15}$NH$_4$Cl as the sole nitrogen source and on uniformly labeled glucose, or acetate or methanol as the carbon source (Batey et al. 1995; Hoffman and Holland 1995; Pardi 1995). RNAs transcribed using these labels inspired new multidimensional NMR experiments that enabled the assignment of chemical shifts to each nucleus in a polynucleotide chain, and made possible the routine structure determination and dynamics characterizations of RNAs 30 nucleotides or less (Latham et al. 2005; Dayie 2008; Lu et al. 2010).

Yet the uniform isotopic labeling methods introduced earlier do not overcome the inherent signal degeneracy problems of large RNAs. Instead, uniform labeling exacerbates the degeneracy problem and negates the benefits of monitoring RNA-ligand interactions, assignment of resonances and structural characterizations in the following fashion. As shown previously, for uniformly labeled nucleotides, the ribose C2' and C4' positions form a triplet that increases the spectral complexity by three-fold. Similarly, the C1' and C5' positions form doublets that increase two-fold the number of resonances in non-constant time HQSC spectra (Dayie and Thakur 2010). Uniform labeling also introduces considerable direct one-bond scalar couplings and residual dipolar couplings from adjacent carbons.

These unwanted splittings can be removed using either constant time experiments (Bax et al. 1979; Bax and Freeman 1981; Grzesiek and Bax 1992; van de Ven and Philippens 1992), adiabatic band selective decoupling schemes (Kupce and Wagner 1996; Brutscher et al. 2001; Dayie 2005), or maximum entropy reconstruction-deconstruction (Shimba et al. 2003). But each method has its attendant disadvantages. Constant time experiments limit the acquisition times to $t_{1}^{\text{max}} = nT_{CC}$ where $n$ is an integer, $T_{CC}$ is the homonuclear coupling constant (e.g. $J_{CC} = 67$ Hz and $J_{C1'/C2'} = 43$ Hz). During the constant-time period, $T$, the decay of the transverse magnetization is proportional to $\exp(-R_2T)$, where $R_2$ is the homogenous transverse rate constant. A compromise must therefore be found between sensitivity (small values of $T$) and resolution (large values of $T$); the long constant-time delays needed to improve resolution lead to significant signal attenuation for RNA molecules larger than 30 nucleotides (Dayie 2005). Additionally, band selective decoupling experiments preclude the observation of the regions to which the decoupling schemes are applied. Finally, the maximum entropy reconstruction-deconstruction requires uniform coupling constants, but in RNA these couplings vary substantially (32–42 Hz in ribose ring).

Selective $^{13}$C labeling schemes can potentially remove most of these drawbacks. By selectively labeling each carbon site, the severe overlap of C2' with C3' is eliminated. In addition, the three-fold complexity created by the triplets formed by C2' and C4' carbons and the doublets formed by the C1' and C5' carbons are removed (Dayie and Thakur 2010). Similarly the direct one-bond scalar couplings and residual dipolar couplings from adjacent carbons introduced by uniform labeling is diminished. By this means the coupling topology is simplified, and signal sensitivity and spectral resolution are enhanced.

These favorable properties make selective $^{13}$C labeling attractive for diverse NMR applications. For example, straightforward measurements of various NMR parameters using standard multidimensional experiments can be envisioned. First, quantitative measurements of relaxation rates of $^{13}$C–$^1$H spin pairs, previously complicated by the presence of $^{13}$C–$^{13}$C dipolar interactions and $^{1}$CC couplings in uniformly labeled samples (Yamazaki et al. 1994; Dayie et al. 2002; Boisbouvier et al. 2003), can now be carried out (Johnson and Hooogstraten 2008) even for large RNA molecules. Second, direct carbon detection experiments suffer from reduction in signal intensity during acquisition because of the evolution of $^{13}$C–$^{13}$C scalar couplings (Bermel et al. 2006; Fiala and Sklenář 2007; Farès et al. 2007). Using selectively labeled samples, these experiments can now be run without complicated pulse programs, or complicated data processing, or both. Third, a number of previously developed methods for measuring residual dipolar couplings and chemical shift anisotropy resulting from weak alignment (Boisbouvier et al. 2004; Miclet et al. 2004; Ying et al. 2006) will benefit considerably using the selective labels. Again straightforward measurements of these couplings can be undertaken without recourse to using filtering elements to remove unwanted $^{13}$C–$^{13}$C J couplings. Finally transverse relaxation optimized spectroscopy (TROSY) studies promise to extend the size of macromolecules that can be studied by NMR (Pervushin et al. 1997). Here again selective labeling can provide a clean and simple alternative for recording TROSY spectra without obliterating the information content of decoupled regions. For example, we expect a two-fold improvement in resolution in the C5' region using methylene TROSY experiments without the need for a constant time evolution and band-selective decoupling, and a two-fold improvement in resolution in the C1' region without sacrificing the information content in the C2' region using non-constant time methine TROSY experiments that dispense with band-selective decoupling.
While alternate $^{13}$C-$^{12}$C labeling schemes have been shown to be important for tackling large protein structural analysis (Religa and Kay 2010; Takeuchi et al. 2010), fewer such studies have been carried out for RNA molecules (Johnson et al. 2006; Hoogstraten and Johnson 2008; Dayie and Thakur 2010) despite some of the potential applications of selective labels enumerated above. For RNA analysis, specific-labeled nucleoside triphosphates (NTPs) are readily synthesized using wildtype and mutant E. coli bacteria as attractive molecular factories. For example, the growth of E. coli K10-1516 strain (glucose-6-phosphate dehydrogenase enzyme ($zwf$ gene) is disabled, hereafter referred to as K10zwf) on $^{13}$C-2-glycerol provides labeling mostly at the C2' and C4' ribose positions, and at both C5 and C6 positions of pyrimidines (Johnson et al. 2006; Hoogstraten and Johnson 2008; Dayie and Thakur 2010). Similarly, growth of an E. coli strain deficient in tricarboxylic acid cycle enzymes malate and succinate dehydrogenases (LeMaster and Kushlan 1996) (DL323) on $^{13}$C-1,3-glycerol yields $^{13}$C label at the C5 position of pyrimidines, and growth on $^{13}$C-2-glycerol yields $^{13}$C label at the C6 position of pyrimidines (Johnson et al. 2006). We recently demonstrated that spiking the growth medium with $^{13}$C labeled formate increases the number of sites selectively enriched in K10zwf (Dayie and Thakur 2010). We observed that labels produced by K10zwf at C1' and C5' are $\sim$7 and $\sim$15%, respectively, whereas the fully enzymatic method can yield $>$90% at those sites. However, the fully enzymatic method is limited to labeling each ribose position using site-specific-labeled glucose that also requires the coupling of the base to the labeled sugar component, a costly undertaking (Tolbert and Williamson 1996; Schultheisz et al. 2008).

Here we explored the potential to preserve a higher level of label at these two important ribose positions and the protonated base sites while retaining the advantage of having an isolated two spin system that will limit the magnetization transfers to desired pathways. The mutant E. coli DL323 strain has the two tricarboxylic acid cycle enzymes succinate and malate dehydrogenases disabled, making it an alternative strain that could compensate for the drawbacks of the K10zwf strain and the enzymatic method. We therefore extended those earlier studies to DL323 and present an analysis of the isotopic enrichment using labeled glycerol and formate in DL323. As expected, we find that it is not necessary to spike the $^{13}$C-1,3-glycerol growth media with $^{13}$C-labeled formate to label the C2 and C8 purine sites. These labeling patterns contrast with the ones obtained using $^{13}$C-2-glycerol wherein exogenous labeled formate is necessary to label the C2 and C8 purine positions. Moreover to explore the usefulness of these site specific labels under conditions of broadened resonances of large RNAs, we dissolved the labeled nucleotides in perdeuterated glycerol and show that CH$_2$-methylene TROSY gives the expected carbon correlations, making this approach likely applicable to high molecular weight RNAs. Obtaining a high degree of labeling at ribose C1', C2', C5' and purine C2 and C8 positions increases the scope of the isotope labeling patterns that are useful for structural, functional and dynamics studies of important RNA macromolecules.

**Materials and methods**

**Bacterial strains**

The mutant strain DL323 (CGSC # 7538, F-, $sdh$-I, $\lambda$lambda $^-$, $mdh$-2, $rph$-I) used in this work was obtained from the Yale Coli Genetic Stock Center.

**Isotopes**

The following isotopically labeled compounds were purchased from Cambridge Isotope Laboratory (Andover, MA) and Isotec-Sigma-Aldrich (Miamisburg, OH): $^{13}$C-sodium formate (99%), $^{13}$C-1,3-glycerol (99%), $^{15}$N-(NH$_4$)$_2$SO$_4$ (99%), and $^{13}$C-2-glycerol (99%).

**Stock solutions**

The stock solutions required for bacterial growth were prepared using distilled and deionized water heat sterilized. The Studier media (SPG) contains 25 mM (NH$_4$)$_2$SO$_4$, 12.6 mM (NH$_4$)$_2$HPO$_4$, 10 mM Na$_2$HPO$_4$, 50 mM KH$_2$PO$_4$, 2 mM MgSO$_4$ and trace metal solution, supplemented with either labeled glycerol or formate and glycerol (Studier 2005). The LeMaster-Richards (LMR) media contains 176 mM KH$_2$PO$_4$, 25 mM NaOH, 10 $\mu$L H$_2$SO$_4$, 12.6 mM (NH$_4$)$_2$SO$_4$, 2 mM MgSO$_4$, 10 $\mu$M FeSO$_4$ and 0.2% trace metals (LeMaster and Richards 1982).

**Media for bacterial growth**

Luria–Bertani (LB), LMR and SPG minimal media were prepared as described (Sambrook and Russell 2001; Studier 2005).

**Growth optimization procedure**

The optimization of mutant DL323 E. coli growth was performed for high yields of biomass per input gram of carbon source (Thakur et al. 2010). Briefly, 5 ml starter cultures in unlabeled SPG or LMR medium were inoculated from a single colony of DL323 grown on LB plates without antibiotics incubated at 37°C overnight. The next
morning, cells were washed twice in 1× phosphate-buffered saline (PBS) and re-suspended in 5 ml of fresh SPG medium without carbon source; 1 ml from this re-suspension was added to 50 ml of SPG medium and incubated at 37°C. After 12–14 h of incubation the cells were washed twice in 1× PBS and resuspended in 50 ml SPG medium with no carbon source, 5 ml from this resuspension were added to 500 ml of SPG medium supplemented with labeled carbon sources and incubated at 37°C for 12 h. Growth in LMR was carried out identically as described above for SPG.

Nucleic acid digestion and boronate column purification

The cell pellet from the 500 ml culture was resuspended in 20 ml lysis buffer (STE buffer: 0.1 M NaCl, 10 mM Tris, 1 mM EDTA, pH 8 and 0.5% SDS). The nucleic acids were separated from the cellular proteins using a 25:24:1 phenol:chloroform:isoamyl alcohol mixture. The nucleic acids present in the upper aqueous layer were extracted multiple times with an equal volume of a 24:1 chloroform:isoamyl alcohol mixture, then the nucleic acids were precipitated overnight at −20°C with ethanol and acetate buffer pH 5.2. The total nucleic acids were subjected to nuclease P1 digestion in 15 mM sodium acetate, 0.1 mM ZnSO4, pH 5.2. The mixture was lyophilized and re-suspended in 10 ml of 1 M triethylammonium bicarbonate (TEABC). Later, the mixture was applied to a cis-diol boronate affinity column to separate deoxyribonucleotides from ribonucleotides. The boronate resin was hydrated in 10 mM Tris HCl, pH 8, rinsed with acidified water, pH 4.5, and equilibrated with 1 M triethylammonium bicarbonate (TEABC), pH 9.5, prior to the separation of the nucleotides. The nucleotide mixture was applied to the column, unbound deoxyribonucleotides and salts were washed with five column volumes of 1 M TEABC, and the unbound deoxyribonucleotides were collected and pooled. The bound ribonucleotides were eluted with acidified water, pH 4.5, and the purity of both dNMPs and rNMPs were checked using NMR.

Separation and desalting of rNMPs by reversed-phase FPLC

The lyophilized rNMPs were dissolved in a minimal amount of solution A (0.1 M monobasic potassium phosphate prepared in distilled water) and passed through a 0.22-µm syringe membrane filter (Fisherbrand) prior to injection on a Targa C18 column (250 × 20 mm, 10 µm, 120 Å, Higgins Analytical, Inc.) using RP-FPLC on an AKTA Design™ system with a detector at a wavelength of 254 nm. Two segment linear gradients of solution B (0.1 M monobasic potassium phosphate, 20% CH3CN, prepared in distilled water) were employed with a flow rate of 10 ml/min and gradients of 0.3 and 10% of buffer B per minute. The sample purity was examined with an analytical RP-FPLC on a Targa C18 column (250 × 4.6 mm, 5 µm, Higgins Analytical, Inc.) with a flow rate of 0.8 ml/min. The purified NMPs were desalted prior to phosphorylation using the same preparative column washed with solution C (0.05% aqueous trifluoroacetic acid (TFA)) and a gradient of 5% solution D (20% CH3CN, 0.05% TFA) per minute with a flow rate of 10 ml/min.

NMR experiments

NMR experiments were run on a four channel Bruker Avance III 600 MHz spectrometer equipped with actively shielded z-axis gradient triple resonance probe at temperatures of 15–45°C. The NMR data sets were processed and the peak positions and intensities were analyzed with Bruker’s TOPSPIN 2.1 as described previously (Dayie and Thakur 2010). One dimensional (1D) 13C spectra and two-dimensional non-constant-time (1H, 13C) heteronuclear single quantum correlation (HSQC) spectra (Bodenhausen and Ruben 1980; Bax et al. 1990) were acquired to analyze the rNMP fractions from the DL323 bacterial strain. To ascertain the labeling pattern of carbon in both the ribose and the base rings, the fractional 13C enrichment at each carbon site was quantified directly by 1D proton methods or indirectly using 2-bond (2JHN) HSQC as described previously (Dayie and Thakur 2010). In addition, the fractional 13C enrichment at each carbon site was quantified by comparing the labeling pattern and intensity of resonances in proton decoupled direct carbon spectra of uniformly 15N-labeled. The fractional enrichment is defined as

\[
\text{Frac}_{i} = \frac{I_{i}}{I_{u}},
\]

where \(I_{i}\) and \(I_{u}\) are the peak intensities of the \(i\)th resonance in the direct 1D or 2D HSQC spectra of the selectively and uniformly labeled 13C/15N rNMP from commercial sources and rNMPs produced with DL323 E. coli. The samples were matched in concentration. However to account for slight differences in sample concentrations, the intensities of each of the spectra were normalized by the average peak intensities in the 2JHN 1HN-15N HSQC data sets because all nucleotides are uniformly 15N-labeled. The fractional enrichment is defined as

\[
\text{Frac}_{i} = \left(\frac{I_{i}}{I_{u}}\right) \left(\frac{N_{i}}{N_{u}}\right),
\]

where \(I_{i}\) and \(I_{u}\) are the peak intensities of the \(i\)th resonance in the direct 1D or 2D HSQC carbon spectra of the selectively and uniformly labeled rNMPs that were recorded identically, \(N_{i}\) and \(N_{u}\) are normalization factors derived from the average peak intensities in the 2JHN 1HN-15N HSQC data sets of the different samples (Lundström et al. 2007). CH2-TROSY experiments (Miclet et al. 2004) were run with the following slight modifications: the WURST-4 decoupling waveform was not applied during the carbon t1 evolution period, and the selective pulse on the CS′ carbon was replaced with a non-selective 180° pulse.
Results and discussion

Incorporation of $^{13}$C into ribose ring of nucleotides using the pentose phosphate pathway (PPP)

To put our results in the context of the *E. coli* mutant DL323 grown in labeled glycerol with or without labeled formate, we present a brief overview of the metabolic pathways for nucleotide metabolism via glycolysis, gluconeogenesis, and the Krebs cycle. Details can be found in standard textbooks (Nelson and Cox 2008; Voet et al. 2008). The ribose moiety of the ribonucleotide is derived exclusively from ribose 5-phosphate (R5P) produced via glycolysis, glucose-6-phosphate (G6P) directly from gluconeogenesis (Thauer et al. 1972; Knappe et al. 1974) possibly by the reversible action of pyruvate formate lyase (Kirkpatrick et al. 2001) (Fig. 1). The purine ring atoms C2 and C8 also derive from the formate pool described above. Thus the purine C2 and C8 sites are expected to be labeled with $^{13}$C using $^{13}$C-1,3-glycerol. If $^{13}$C-1,3-glycerol is used addition of exogenous formate should make no difference to labeling at the purine C2 and C8 positions.

For DL323 *E. coli* grown in $^{13}$C-1,3-glycerol without labeled formate, the ribose ring is labeled in all but the C4' carbon position (Fig. 2) in agreement with metabolic analysis (Fig. 1) and previous work (Johnson et al. 2006). As a result, the C2' and C3' positions suffer from splitting and condenses it with E4P to form sedoheptulose 7-phosphate (S7P) and GA3P. In the third reverse reaction, transaldolase (TAL) removes a three carbon fragment from F6P and condenses it with E4P to form sedoheptulose 7-phosphate (S7P) and GA3P. In the third reverse reaction, transketolase (TKL) catalyzes the transfer of a two carbon fragment from the ketose donor F6P to the aldose acceptor GA3P to form xylulose 5-phosphate (X5P) and erythrose 4-phosphate (E4P). This X5P, eventually isomerized to R5P, is labeled at only the C2 and C4 positions. In the second reverse reaction, transaldolase (TAL) removes a three carbon fragment from F6P and condenses it with E4P to form sedoheptulose 7-phosphate (S7P) and GA3P. In the third reverse reaction, transketolase again transfers a two carbon fragment from S7P to GA3P to form X5P and R5P. In this case, X5P is labeled at C2 and C4 whereas R5P is labeled at only C4. The predicted net effect is a 2:1 labeling pattern of [2, 4-$^{13}$C]ribose and [4-$^{13}$C]ribose, and no label at the 3 or 5 positions via the noPPP. The oPPP will lead to a [1, 4-$^{13}$C] ribose label (Fig. 1) and unlabeled carbon dioxide. A similar analysis indicates that placement of the $^{13}$C-label at the terminal C-1,3 carbons of glycerol leads to the following labeling pattern. The reverse noPPP leads a 2:1 labeling pattern of [1, 3, 5-$^{13}$C] ribose and [1, 2, 3, 5-$^{13}$C] ribose with no label at the 4 position. The oPPP will lead to a [2, 3, 5-$^{13}$C] ribose label, but no label at the 4-position (Fig. 1). Thus one expects coupling between ribose C1' and C2', and between C2' and C1' and C3'.

Incorporation of $^{13}$C into base ring of nucleotides via the glycolysis, gluconeogenesis and the Krebs cycle

The main advantage of the DL323 strain is that the flux through the TCA cycle is reduced to zero thereby preventing the dilution of the labels arising from oxaloacetate (LeMaster and Kushlan 1996). The nucleotide bases are synthesized from the precursors of 3PG that give rise to glycine (Gly) and serine (Ser) and from oxaloacetate that gives rise to aspartic acid (Asp). The amide group, the C2'-, C6'- and C7'-carbon positions of Asp become the N1, C6, C5, and C4 ring atoms, respectively of pyrimidines (Fig. 1). The bicarbonate pool arises from several sources: the decarboxylation of PEP to form acetyl-CoA, carbon 1 of G6P and carbon 1 of pyruvate (derived from carbon 1 or 3 of glycerol). The formate pool also arises from the C6H2OH of Ser (derived from carbon 1 or 3 of glycerol). In addition, exogenous formate could enter the metabolic cycle by exchanging the carbonyl group of pyruvate by consuming acetyl-CoA (Thauer et al. 1972; Knappe et al. 1974) possibly by the reversible action of pyruvate formate lyase (Kirkpatrick et al. 2001) (Fig. 1). The purine ring atoms C2 and C8 also derive from the formate pool described above. Thus the purine C2 and C8 sites are expected to be labeled with $^{13}$C using $^{13}$C-1,3-glycerol. If $^{13}$C-1,3-glycerol is used addition of exogenous formate should make no difference to labeling at the purine C2 and C8 positions.

For DL323 *E. coli* grown in $^{13}$C-1,3-glycerol without labeled formate, the ribose ring is labeled in all but the C4' carbon position (Fig. 2) in agreement with metabolic analysis (Fig. 1) and previous work (Johnson et al. 2006). As a result, the C2' and C3' positions suffer from splitting and condenses it with E4P to form sedoheptulose 7-phosphate (S7P) and GA3P. In the third reverse reaction, transaldolase (TAL) removes a three carbon fragment from F6P and condenses it with E4P to form sedoheptulose 7-phosphate (S7P) and GA3P. In the third reverse reaction, transketolase (TKL) catalyzes the transfer of a two carbon fragment from the ketose donor F6P to the aldose acceptor GA3P to form xylulose 5-phosphate (X5P) and erythrose 4-phosphate (E4P). This X5P, eventually isomerized to R5P, is labeled at only the C2 and C4 positions. In the second reverse reaction, transaldolase (TAL) removes a three carbon fragment from F6P and condenses it with E4P to form sedoheptulose 7-phosphate (S7P) and GA3P. In the third reverse reaction, transketolase again transfers a two carbon fragment from S7P to GA3P to form X5P and R5P. In this case, X5P is labeled at C2 and C4 whereas R5P is labeled at only C4. The predicted net effect is a 2:1 labeling pattern of [2, 4-$^{13}$C] ribose and [4-$^{13}$C] ribose, and no label at the 3 or 5 positions via the noPPP. The oPPP will lead to a [1, 4-$^{13}$C] ribose label (Fig. 1) and unlabeled carbon dioxide. A similar analysis indicates that placement of the $^{13}$C-label at the terminal C-1,3 carbons of glycerol leads to the following labeling pattern. The reverse noPPP leads a 2:1 labeling pattern of [1, 3, 5-$^{13}$C] ribose and [1, 2, 3, 5-$^{13}$C] ribose with no label at the 4 position. The oPPP will lead to a [2, 3, 5-$^{13}$C] ribose label, but no label at the 4-position (Fig. 1). Thus one expects coupling between ribose C1' and C2', and between C2' and C1' and C3'.

For DL323 *E. coli* grown in $^{13}$C-2-glycerol without labeled formate, the ribose ring is labeled in all but the C3' and C5' carbon positions (Fig. 3) in agreement with metabolic analysis (Fig. 1) and previous work (Johnson et al. 2006). As a result, the C2' and C4' positions are not split by the carbon–carbon coupling observed in strain DL323 grown in $^{13}$C-1,3-glycerol. The C1' positions also remain...
Fig. 1 Major metabolic pathways involved in the production of nucleic acid nucleotides from glycerol, including key steps in glycolysis, gluconeogenesis and one pass through the tricarboxylic (TCA) cycle. With the E. coli strain lacking succinate and malate dehydrogenase (DL323), the oxidative branch of the pentose phosphate pathway remains intact but the TCA cycle is severed in two places such that the oxaloacetate is derived exclusively from carboxylation of PEP and the resulting label is not diluted by the TCA cycle. Atom labels for the terminal (1,3) carbons (magenta and thin circle) and central (2) carbon (cyan and thick circle) of glycerol are highlighted. Positions that are enriched due to the presence of $^{13}$CO$_2$ in the growth medium are shown with an X. Pyrimidine base derived from the Oxaloacetate (OAA) produced by carboxylation of phosphoenolpyruvate (PEP) is shown via the aspartate intermediate. This OAA cannot be used as a substrate in the first and subsequent rounds of the TCA cycle because of the two mutations. Consequently OAA derived aspartate amino acid can be produced with $^{13}$C labeling at only the C$^a$ position if $^{13}$C-2-glycerol is used. If $^{13}$C-1,3-glycerol is used only C$^b$ position is labeled. In either case carboxylation of PEP leads to labeling of the C$^c$ position. Similarly the labeling pattern of purines from glycine derived from 3PG are shown such that if $^{13}$C-2-glycerol is used only the C$^a$ position of Gly and therefore C5 position of the purine ring is labeled. Otherwise if $^{13}$C-1,3-glycerol is used the CO of Gly and therefore C4 of purine ring is labeled, and the C$^b$ position of Ser leads to labeling of the purine C2 and C8 positions.
singlet (Fig. 3), presumably the labeled C2 arises in a different molecule (Fig. 1). As expected, only the pyrimidine C6 base atoms are substantially labeled to \( \approx 96\% \). The C2 and C8 atoms of purine are labeled to \( \approx 5\% \), comparable to that observed for K10zwf E. coli grown in \(^{13}\)C-1,3-glycerol without labeled formate (Dayie and Thakur 2010).

In contrast to K10zwf E. coli grown in \(^{13}\)C-2-glycerol with labeled formate (Dayie and Thakur 2010), addition of labeled formate to the \(^{13}\)C-2-glycerol media does not lead to new labels at the C3 and C5 positions (Fig. 3c). Similar to K10zwf E. coli, addition of labeled formate to \(^{13}\)C-2-glycerol leads to increased labeling of the C8 carbon positions of the purine ring to \( \approx 88\% \) (Fig. 3). Surprisingly, C2 remains labeled at low levels unlike the case seen in the K10zwf strain (Dayie and Thakur 2010).

### Estimating the level of carbon labeling efficiency

The level of \(^{13}\)C labeling efficiency is easily estimated using 1D \(^1\)H or natural abundance \(^{13}\)C carbon spectra. However, for cases of overlap, two other methods can be used. The first method exploits long range proton-nitrogen correlations in \(^{15}\)N-HSQC spectra to estimate the labeling efficiency of the purine C2 and C8 carbon sites, the pyrimidine C5 and C6 carbon sites and the purine C1 carbon site (Fig. 4). The relaxation properties and transfer efficiencies are different for long range and one-bond magnetization transfers; nonetheless, the use of the long range \(^{15}\)N-HSQC method has been shown to be useful for estimating the level of \(^{13}\)C incorporation (Dayie and Thakur 2010). The labeling efficiency is estimated by...
comparing the intensity of each $^{13}$C satellite peak to the intensity of the center peak. For a mixture of the four rNMPs extracted from DL323 E. coli grown in $^{13}$C-1,3-glycerol with or without labeled formate, there is significant overlap in both the base and ribose regions even for an isolated rUMP (Fig. 2). Analysis of the 2D $^2$J$_{HN}$ HSQC spectra (Fig. 4) allows a straightforward estimate of the labeling at purine C2 and C8 and pyrimidine C6 (Fig. 4). For site-specific $^{13}$C/$^{15}$N-labeled rNMP, the 2D method is in excellent agreement with the 1D $^1$H method (Fig. 4). In the absence of formate, the level of enrichment was ~90% for the purine C8 and C2. In the presence of formate the level of enrichment remains at ~90% for purine C8 and ~90% for Ade C2 (Table 1). As expected, each of the H2 and H8 proton resonance is split into a doublet with little central peak in the acquisition dimension (Fig. 4a). In contrast, each H6 proton resonance gives rise to a singlet suggestive of almost no label at the pyrimidine C6 position (Fig. 4a); each H5 proton resonance shows a doublet suggestive of almost complete labeling (not shown). The C1' and C5' regions were sufficiently isolated to enable an estimate of the labeling efficiencies using the 1D $^1$H NMR spectrum: the enrichment level was ~58% for the ribose C1' and 90% for the ribose C5'.

Similarly, analyses of the multiplet pattern of the four labeled nucleotides derived from DL323 E. coli grown in $^{13}$C-2-glycerol without formate show the opposite labeling pattern in the purine C2/C8 and pyrimidine C5/C6 regions. As expected, the H2 proton resonance gives rise to a singlet (at N1 and N3) suggestive of little labeling at the purine C2...
Table 1 13C enrichment levels at various carbon positions within ribonucleotides harvested from DL323 E. coli grown on 13C-1,3-glycerol with and without 13C-formate as carbon sources

| Purine position | 13C-Carbon source: 1,3-Glycerol only | 13C-Carbon source: 1,3-Glycerol and Formate |
|-----------------|-------------------------------------|--------------------------------------------|
| Ade C2          | 0.88 ± 0.04                         | 0.82 ± 0.13                                |
| C8              | 0.92 ± 0.07                         | 0.81 ± 0.13b                               |
| Pyrimidine C5   | 0.88 ± 0.04                         | 0.92 ± 0.03                                |
| C6              | <0.05                               | 0.22 ± 0.02                                |
| Ribose C1      | 0.58 ± 0.08                         | 0.47 ± 0.11                                |
| C2             | 0.50 ± 0.04                         | 0.57 ± 0.09                                |
| C3             | 0.93 ± 0.01                         | 0.77 ± 0.07                                |
| C4             | <0.02                              | <0.02                                      |
| C5             | 0.94 ± 0.03                         | 0.85 ± 0.06                                |

a The percentage label is calculated as an average of three methods: (i) the ratio of the sum of the intensities of satellite peaks to the sum of the intensities of the satellite and center peaks using the 2-bond 15N HSQC without 13C decoupling during acquisition (Dayie and Thakur 2010); (ii) the ratio of the sum of the intensities of satellite peaks to the sum of the intensities of the satellite and center peaks using the 1D 1H experiment without 13C decoupling during acquisition and (iii) using the fractional enrichment (Frac_E) method of Lundström et al. (2007) as described in the text. We find that direct 1D and 2D methods (i and ii) give consistently slightly higher values than the fractional enrichment method (iii). 

b The percentage label (Plabel) is calculated as in (a) but this time with only method (iii) 

The labeling pattern of rNMPs derived from DL323 grown in 13C-1,3-glycerol and formate is the case for growth in 13C-1,3-glycerol, gives rise to a doublet with little central peak in the acquisition dimension (Fig. 4a) indicating substantial labeling (~98%) at the pyrimidine C6 position (Fig. 4c). This method using the long range proton-nitrogen correlations in 15N-HSQC spectra is adequate for estimating the labeling efficiency in the aromatic ring (Tables 1, 2). The method is, however, limited to purine C2 and C8 carbon sites, the pyrimidine C5 and C6 carbon sites and the purine C1’ carbon site. The second method of fractional enrichment (Lundström et al. 2007) gets around this limitation. However this fractional enrichment method cannot be carried out on a mixture of the four nucleotides obtained from boronate affinity purification. The four nucleotides must first be separated. Then each component can be analyzed accurately. Using this method we obtain enrichment values that are comparable to but consistently larger (by about 5–30%) than those obtained with the two previous methods. The values reported in Tables 1 and 2 are the averages of the three methods.

Applications of selective labels for 13C NMR study of nucleic acids

An example of the high quality uncluttered spectra obtained using strain DL323 is shown in Fig. 5. The labeling pattern of rNMPs derived from DL323 grown in 13C-2-glycerol and formate is particularly attractive for relaxation studies of not only the ribose C2’ and C4’ positions, as elegantly demonstrated recently by Hoogstraten and his colleagues (Johnson et al. 2006; Johnson and Hoogstraten 2008), but also the ribose C1’ and the base C6 and C8 positions. Each of these five positions is essentially singlet. Their isolation from directly labeled adjacent neighbors means that the interference arising from strong 13C–13C magnetic interactions in the base and ribose rings is no longer an issue for extracting accurate relaxation parameters (Johnson et al. 2006; Johnson and Hoogstraten 2008).

The labeling pattern of rNMPs derived from DL323 grown in 13C-1,3-glycerol and formate is less attractive for relaxation studies of the ribose region because the C2’ position is doublet, the C1’ retains some residual doublet arising from 13C2−13C1’ isotopomers, and the C4’ has no label (Fig. 5a, c). This is consistent with the metabolic analysis that predicts that at most 20% of the molecules have both C1’ and C2’ labeled, whereas 75% of the molecules have either C2’ and C1’ labeled or C2’ and C3’ labeled (Fig. 1). In principle, measurements on isolated 13C1’ spins can be performed so long as suitable purging pulse sequences are employed to remove 13C2’−13C1’ pairs. In the base region only the purine C2 and C8 sites are available for clean relaxation measurements; the pyrimidine C5 site has multiplet structure precluding its use for accurate relaxation measurements.

Nonetheless, an attractive feature of the labels derived from DL323 grown in 13C-1,3-glycerol and formate is the...
complete isolation of the C5' labels from C4'. This label is therefore appropriate for NMR applications that exploit the methylene carbon position in RNA. As demonstrated by Bax and coworkers, considerable gain in 1H resolution is possible for methylene C5' regions in a 19-base-pair DNA fragment (Miclet et al. 2004).

Therefore, to ascertain the usefulness of site specific labels under conditions of broadened resonances that accompany RNA of increased size, we dissolved the labeled nucleotides in perdeuterated glycerol. At a temperature of 40°C, the ribose C5' resonances are not only broadened in a normal non-constant time 13C HSQC experiment (Fig. 6c) but also exhibit extensive spectral overlap. To estimate the overall tumbling time constant (τc), at this temperature, we use the high frequency approximation (Farrow et al. 1995; Ishima and Nagayama 1995; Peng and Wagner 1995; Dayie et al. 1996) and assume isotropic tumbling, such that the overall correlation time is independent of the order parameter (Kay et al. 1989; Eldho and Dayie 2007):

$$\tau_c \approx \left( \frac{\sqrt{6R_2/R_1 - 7}}{2\omega_X} \right)$$

where $R_1$ is the longitudinal relaxation rate constant, $R_2$ is the transverse relaxation rate constant, and $\omega_X$ is the Larmor frequency of the X (15N or 13C) spin. At 40°C, we estimate the rotational correlation time to be 9.5 ± 0.7 ns. Given 6 ns as the correlation time for a 36 nucleotide RNA at
Table 2  

| Carbon position labeled | 13C-Carbon source: 2-Glycerol only | 13C-Carbon source: 2-Glycerol and Formate |
|-------------------------|-----------------------------------|------------------------------------------|
| Purine                  |                                   |                                          |
| Ade C2                  | 0.16 ± 0.10                       | 0.12 ± 0.06                              |
| C8                      | 0.17 ± 0.12                       | 0.88 ± 0.03                              |
| Pyrimidine              |                                   |                                          |
| C5                      | <0.03                             | <0.03                                    |
| C6                      | 0.98 ± 0.003                      | 0.95 ± 0.02                              |
| Ribose                  |                                   |                                          |
| C1'                     | 0.51 ± 0.11                       | 0.33 ± 0.05                              |
| C2'                     | 0.30 ± 0.09                       | 0.60 ± 0.08                              |
| C3'                     | <0.02                             | <0.02                                    |
| C4'                     | 0.87 ± 0.22                       | 1.0 ± 0.17                               |
| C5'                     | <0.02                             | <0.02                                    |

* The percentage labels are calculated as described in the footnote to Table 1.

25°C (Eldho and Dayie 2007), and assuming a linear correlation between molecular weight and correlation time, then these nucleotides are behaving roughly as a 50–60 nucleotide RNA. Experiments carried out at lower temperatures (down to 15°C) indicate useful signals can be obtained for RNAs of up to 150 nucleotides (~30 ns correlation time) in agreement with previous observations (Hansen and Al-Hashimi 2007).

Use of the non-constant time version of the 13CH2 TROSY experiment (Miclet et al. 2004) enables the rescue
of the slowest relaxing multiplet component with improved resolution and sensitivity compared to the fastest relaxing and broadened signals (Fig. 6a, b). Compared to the normal CH$_2$-methylene optimized HSQC (Schleucher et al. 1994; Sattler et al. 1995), all four rNMPs in the mixture are resolved at 800 MHz proton frequency. It is anticipated that these and other new experiments that incorporate the $^{13}$CH$_2$ TROSY module can be designed to probe RNA-ligand interactions at very high resolution using the site specific labels described here.

As discussed previously for the K10zwf labeled rNMPs (Dayie and Thakur 2010), the DL323 labeled rNMPs will also benefit a number of important spin relaxation applications. For instance, obtaining accurate relaxation parameters from $^{13}$C-CPMG-based relaxation dispersion experiments for quantifying millisecond time-scale processes is predicated on removing modulations that complicate analysis of the data (Yamazaki et al. 1994; Johnson et al. 2006; Johnson and Hoogstraten 2008; Hansen et al. 2009).

**Conclusion**

*Escherichia coli* strains with lesions in specific metabolic pathways are attractive for synthesizing selectively labeled nucleotides for tackling large RNA macromolecules (Johnson et al. 2006; Johnson and Hoogstraten 2008; Dayie and Thakur 2010). Here we show that we can preserve a high level of label at the ribose C1$'$ and C5$'$ and base C2 and C8 without introducing labels in adjacent carbon positions, thereby ensuring an isolated two spin system that limits extraneous relaxation pathways. The use of the mutant *E. coli* DL323 strain that is deficient in succinate and malate dehydrogenases has the added advantage that the flux through the TCA cycle is reduced to zero such that maximal labeling is obtained at either the C5 or C6 pyrimidine position without compromising the label of the ribose ring. The use of $^{13}$C-1,3-glycerol produces rNMPs enriched at C1$'$ ($\sim$ 60%) and C5$'$ ($\sim$ 90%) in the ribose ring and C2 ($\sim$ 90%) and C8 ($\sim$ 90%) in the purine ring without the deleterious one-bond $^{13}$C–$^{13}$C scalar couplings that could interfere with accurate spin-relaxation measurements. The use of $^{13}$C-2-glycerol also produces rNMPs with enrichment of $^{13}$C at C2$'$ and C4$'$ of the ribose groups and C6 groups in the Pyr ring without incorporation of label at either base C5 sites or ribose C3$'$ or C5$'$. Thus the one-bond $^{13}$C–$^{13}$C scalar couplings that could interfere with accurate spin-relaxation measurements are absent. As expected, we find that the C2 and C8 purine sites are highly $^{13}$C-enriched in rNMPs derived from DL323 strain grown in $^{13}$C-1,3-glycerol growth media without $^{13}$C-labeled formate; rNMPs derived from DL323 strain grown in $^{13}$C-2-glycerol growth media require exogenous formate for high enrichment of mostly the C8 positions. The isolated but highly enriched C5$'$ position opens up an attractive avenue for CH$_2$-methylene TROSY applications for large RNA molecules. Overall then, these labeling patterns of the slowest relaxing multiplet component with improved resolution and sensitivity compared to the fastest relaxing and broadened signals (Fig. 6a, b). Compared to the normal CH$_2$-methylene optimized HSQC (Schleucher et al. 1994; Sattler et al. 1995), all four rNMPs in the mixture are resolved at 800 MHz proton frequency. It is anticipated that these and other new experiments that incorporate the $^{13}$CH$_2$ TROSY module can be designed to probe RNA-ligand interactions at very high resolution using the site specific labels described here.

As discussed previously for the K10zwf labeled rNMPs (Dayie and Thakur 2010), the DL323 labeled rNMPs will also benefit a number of important spin relaxation applications. For instance, obtaining accurate relaxation parameters from $^{13}$C-CPMG-based relaxation dispersion experiments for quantifying millisecond time-scale processes is predicated on removing modulations that complicate analysis of the data (Yamazaki et al. 1994; Johnson et al. 2006; Johnson and Hoogstraten 2008; Hansen et al. 2009).

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![Fig. 6](image-url)
provide a large number of ‘isolated’ backbone ribose and base side-chain probes that are available for the study of dynamics and structure of large RNAs.

Acknowledgments This work was supported in part by the University of Maryland Nano-Biotechnology Award and the National Institutes of Health grant GM077326 to T. K. Dayie, and J. N. Sama is a recipient of a Percy Julian Fellowship (NSF grant CHE0746446) and an HHMI undergraduate fellowship.

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