A Cyclic Metabolic Network in *Pseudomonas protegens* Pf-5 Prioritizes the Entner-Doudoroff Pathway and Exhibits Substrate Hierarchy during Carbohydrate Co-Utilization

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ABSTRACT The genetic characterization of *Pseudomonas protegens* Pf-5 was recently completed. However, the inferred metabolic network structure has not yet been evaluated experimentally. Here, we employed 13C-tracers and quantitative flux analysis to investigate the intracellular network for carbohydrate metabolism. In lieu of the direct phosphorylation of glucose by glucose kinase, glucose catabolism was characterized primarily by the oxidation of glucose to gluconate and 2-ketogluconate before the phosphorylation of these metabolites to feed the Entner-Doudoroff (ED) pathway. In the absence of phosphofructokinase activity, a cyclic flux from the ED pathway to the upper Embden-Meyerhof-Parnas (EMP) pathway was responsible for routing glucose-derived carbons to the non-oxidative pentose phosphate (PP) pathway. Consistent with the lack of annotated genes in *P. protegens* Pf-5 for the transport or initial catabolism of pentoses and galactose, only glucose was assimilated into intracellular metabolites in the presence of xylose, arabinose, or galactose. However, when glucose was fed simultaneously with fructose or mannose, co-uptake of these hexoses was evident, but glucose was preferred over fructose (3 to 1) and over mannose (4 to 1). Despite gene annotation of mannose catabolism to fructose-6-phosphate, metabolite labeling patterns revealed that mannose was assimilated into fructose-1,6-bisphosphate, similarly to fructose catabolism. Remarkably, carbons from mannose and fructose were also found to cycle backward through the upper EMP pathway toward the ED pathway. Therefore, the operational metabolic network for processing carbohydrates in *P. protegens* Pf-5 prioritizes flux through the ED pathway to channel carbons to EMP, PP, and downstream pathways.

IMPORTANCE Species of the *Pseudomonas* genus thrive in various nutritional environments and have strong biocatalytic potential due to their diverse metabolic capabilities. Carbohydrate substrates are ubiquitous both in environmental matrices and in feedstocks for engineered bioconversion. Here, we investigated the metabolic network for carbohydrate metabolism in *Pseudomonas protegens* Pf-5. Metabolic flux quantitation revealed the relative involvement of different catabolic routes in channeling carbohydrate carbons through a cyclic metabolic network. We also uncovered that mannose catabolism was similar to fructose catabolism, despite the annotation of a different pathway in the genome. Elucidation of the constitutive metabolic network in *P. protegens* is important for understanding its innate carbohydrate processing, thus laying the foundation for targeting metabolic engineering of this untapped *Pseudomonas* species.

KEYWORDS *Pseudomonas*, carbohydrate co-utilization, hexose sugar, metabolic flux analysis, metabolomics

Species of the genus *Pseudomonas*, which are ubiquitous in the environment, are metabolically diverse and often touted for industrial bioproduction (1). The elucidation of the cellular network of carbon fluxes through metabolic pathways of these
bacterial species is critical both to understand their role in carbon processing in environmental matrices and to engineer these species to optimize their use in agriculture, industry, and medicine. Gaining importance in bioremediation, _Pseudomonas protegens_ Pf-5 was identified to produce enzymes that degrade polyurethane, a plastic polymer (2). Furthermore, _P. protegens_ Pf-5 is also known to synthesize and release several antimicrobials and exoenzymes that are toxic to plant pathogens (3–6). Recently, _P. protegens_ Pf-5 was characterized and annotated at the genomic level (7). However, the metabolic network of _P. protegens_ Pf-5 has only been inferred from the genome annotation and has not yet been investigated experimentally.

Given the importance and ubiquity of carbohydrate-containing feedstocks, we sought to unravel the metabolic network structure for carbohydrate metabolism in _P. protegens_ Pf-5 by combining $^{13}$C-assisted cellular carbon mapping with $^{13}$C metabolic flux analysis (MFA). Previous studies on other _Pseudomonas_ species (i.e., _P. putida_ and _P. fluorescens_) focused on elucidating the metabolic fluxes during growth on glucose, a prototypical carbohydrate substrate (8–11). In a similar fashion, we also studied the innate carbohydrate metabolism in _P. protegens_ Pf-5 during feeding on glucose alone. However, carbon feedstocks are typically composed of other carbohydrates in addition to glucose. Therefore, we also investigated carbon assimilation and fluxes when the _P. protegens_ Pf-5 cells were fed on mixtures of glucose with other hexoses (mannose, fructose, and galactose) or pentoses (xylose and arabinose).

Previous reports showed that _P. protegens_ strains were able to grow on glucose, mannose, or fructose as a single carbon source but not on galactose, xylose, or arabinose (6). In the genome of _P. protegens_ Pf-5, genes for the following transporters were found: a phosphoenolpyruvate (PEP)/sugar phosphotransferase system (PTS) for fructose uptake (and possibly for mannose uptake) and an ATP-binding cassette (ABC) transporters for glucose, mannose, galactose, and xylose (Fig. 1; see also Table S1 in the supplemental material) (7). Characteristically in _Pseudomonas_ species, glucose metabolism involves a peripheral pathway in the periplasm wherein glucose dehydrogenase converts glucose to gluconate and gluconate 2-dehydrogenase converts gluconate to 2-ketogluconate (Fig. 1) (8). Fluxes through these oxidation reactions were found to maximize growth on glucose (8). After active transport into the cytoplasm, the oxidized products of glucose are phosphorylated to 6-phosphogluconate (6P-gluconate) and subsequently routed to the Entner-Doudoroff (ED) pathway or the pentose phosphate (PP) pathway (Fig. 1) (8, 10, 11). Previous studies with other _Pseudomonas_ species have determined that the forward Embden-Meyerhof-Parnas (EMP) pathway was not possible due to the absence of 6-phosphofructokinase to convert fructose-6-phosphate (F6P) to fructose-1,6-bisphosphate (FBP) (Fig. 1) (1, 7, 10). Therefore, to route glucose-derived carbons eventually toward biosynthetic pathways and the tricarboxylic acid (TCA) cycle, the ED pathway is required, wherein 6P-gluconate is cleaved to produce pyruvate and glyceraldehyde 3-phosphate (GAP) (Fig. 1). The gene that encodes 6-phosphofructokinase was also found to be absent in _P. protegens_ Pf-5 (7); the ED pathway is thus assumed to be the required route for glucose metabolism in _P. protegens_ Pf-5. However, the extent to which the peripheral pathway of glucose oxidation contributes to the initial glucose catabolism compared to direct glucose phosphorylation to glucose-6-phosphate (G6P) in _P. protegens_ Pf-5 remains to be determined.

In contrast to glucose, fructose is transported through the PTS, which uses the phosphate group from PEP to phosphorylate fructose to fructose-1-phosphate (F1P) followed by a subsequent phosphorylation step by 1-phosphofructokinase to convert F1P to FBP (Fig. 1) (9, 12, 13). Previous studies on other _Pseudomonas_ species (_P. fluorescens_, _P. putida_, _P. aeruginosa_, _P. stutzeri_, and _P. acidovorans_) found that fructose-derived carbons were cycled backward via a reverse (gluconeogenic) flux through the upper EMP pathway (FBP up to G6P) to be connected to the ED pathway (12, 14–16). However, according to the genome of _P. protegens_ Pf-5, FBP could also be involved in a forward flux of the EMP pathway, wherein FBP is lysed directly to glyceraldehyde-3-phosphate (GAP) and dihydroxyacetone-3-phosphate (DHAP) (Fig. 1) (7). Whether the
preferential route for fructose assimilation during growth on mixed carbohydrates is via the direct route of FBP to the triose phosphates or through the reverse cycling of carbons from FBP to the ED pathway remains to be determined. The catabolic routing of FBP has important energetic implications for \textit{P. protegens} Pf-5. Compared to the direct FBP lysis through the forward EMP pathway, the combination of reverse flux through the upper EMP pathway with the ED pathway maintains the same quantity of reduced equivalents [i.e., NAD(P)H] but half the ATP yield.

With respect to initial mannose catabolism, the authors of previous studies with \textit{P. aeruginosa} proposed two possible routes, which involve either mannose isomerization to fructose followed by subsequent phosphorylation to FBP or direct phosphorylation of mannose to mannose-6-phosphate (M6P) prior to isomerization to F6P (17, 18). Relevant to the first route, an intracellular mannose isomerase (EC 5.3.1.7) was reported in \textit{P. cepacia} and \textit{P. saccharophila} (19, 20), but the gene for this enzyme was not annotated in the \textit{P. protegens} Pf-5 genome (7). On the other hand, albeit not yet confirmed by metabolic studies, the genes for the relevant enzymes in the second catabolic route, i.e., the conversion of mannose to F6P, were annotated in the \textit{P. protegens} Pf-5 genome (Fig. 1).
Both oxidative and non-oxidative routes of the PP pathway are important to channel carbohydrate-derived metabolite precursors to the biosynthesis of ribonucleotides and aromatic amino acids. The presence of the annotated genes for transketolase and transaldolase enzymes implied a fully functional PP pathway in *P. protegens* Pf-5 (7). For xylose catabolism, the annotation of a ribose transporter indicates that it could be used as a possible xylose transporter, but the genes encoding the enzymes (xylose isomerase and xylulose kinase) responsible for introducing xylose into the PP pathway were not present in *P. protegens* Pf-5 (Fig. 1) (7). Moreover, the collective enzymes needed for the alternative route for xylose through the Weimberg pathway, which incorporates xylose through xylonate into α-ketoglutarate, were not all present in the genome of *P. protegens* Pf-5 (7). Regarding arabinose catabolism, there was no annotated pathway for the assimilation of arabinose in *P. protegens* Pf-5 (Fig. 1) (7). Despite the lack of the relevant genes for xylose catabolism, a recent study found extracellular xylose depletion by *P. protegens* Pf-5 during growth on a mixture of carbohydrates (21). Whether arabinose or xylose is incorporated into the PP pathway by some other non-annotated pathway in the presence of another carbohydrate remains to be determined.

Here we applied liquid chromatography (LC) with high-resolution mass spectrometry (HRMS) to perform a 13C-assisted metabolomics investigation during growth of *P. protegens* Pf-5 on glucose alone or simultaneously with fructose, mannose, galactose, xylose, or arabinose. We provide the first quantitative evaluation of the hypothetical metabolic network of *P. protegens* Pf-5 deduced from its genome-level characterization.

**RESULTS**

**Physiological parameters of *P. protegens* Pf-5 grown on carbohydrate mixtures.** We investigated the growth phenotype of *P. protegens* during batch culturing on different hexose combinations. Our growth rates were in close agreement with the reported values for glucose-grown *P. putida* (0.56 ± 0.02 h⁻¹) and *P. fluorescens* (0.49 ± 0.03 h⁻¹) (8, 22). Starting with a total carbon concentration of 100 mM in the growth medium, the growth rate of cells fed on glucose alone (0.56 ± 0.09 h⁻¹) was similar to the growth rate of cells fed on a 1:1 mixture of glucose and fructose (0.52 ± 0.06 h⁻¹) or glucose and mannose (0.50 ± 0.04 h⁻¹) (Fig. 2A; see also Fig. S1 and Table S2 in the supplemental material). We also found that the growth rate remained relatively unchanged when the cells were grown on 50 mM C glucose alone (0.47 ± 0.02 h⁻¹) (Fig. 2A). Therefore, *P. protegens* would not be subjected to carbon limitation during growth on carbohydrate mixtures if either fructose or mannose was not assimilated from the mixture with glucose (Fig. 2A).

We monitored substrate consumption by the *P. protegens* Pf-5 cells by tracking the depletion of the carbohydrates from the extracellular medium (Fig. 2B; Table S2). The glucose-grown cells depleted glucose completely over a 10-h period (Fig. 2B). Over the same time period, the cells grown on the mixture of glucose and mannose depleted both substrates completely, but cells grown on the mixture of glucose and fructose depleted glucose completely and approximately 70% of fructose (Fig. 2B). We also observed that, during the growth of *P. protegens* Pf-5 on the mixture of glucose and mannose, fructose appeared in the extracellular medium approximately 1 h after mannose consumption started (Fig. 2A). The appearance of extracellular fructose implied the presence of a mannose isomerase, which is responsible for converting mannose to fructose in other *Pseudomonas* species (19, 20) (Fig. 2B). During growth on
both hexose mixtures, glucose was consumed faster and depleted by 6 h of growth, at which time approximately 30% of the fructose was consumed but only 10% of the mannose was depleted by the cells (Fig. 2B). Thus, during hexose co-utilization under both of our experimental conditions, glucose provided the major carbon source for cellular metabolism.

We also monitored the extracellular overflow of metabolic products, a phenomenon that is widely reported in Pseudomonas species (8, 11, 23). Both oxidized products of glucose (i.e., gluconate and 2-ketogluconate) and the organic acid pyruvate were found at appreciable levels (>1 μM) in the extracellular medium (Fig. 2C). Secretions of gluconate and 2-ketogluconate were reported previously with P. putida (8, 11) and P. fluorescens (23); pyruvate secretion was reported previously with P. fluorescens (23). We found that the level of these secreted metabolites from P. protegens Pf-5 was dependent on the substrate composition in the growth medium. The highest secretions of gluconate and 2-ketogluconate (greater than 2 mM) were measured in the medium
when cells were grown on glucose alone or glucose with fructose. In contrast, during growth on the mixture glucose and mannose, the highest secretion of gluconate and 2-ketogluconate decreased substantially, by ~40% and ~85%, respectively (Fig. 2C; Table S2). Compared to gluconate and 2-ketogluconate, pyruvate was secreted in smaller amounts (micromolar range), with the highest secretion (0.13 ± 0.08 mM) obtained when cells were grown on glucose alone (Fig. 2C; Table S2).

**Metabolic pathways and fluxes in glucose-fed cells.** Isotopic enrichment with [1,2-13C2]glucose was used to determine the metabolic network structure through the initial glucose catabolism, the EMP pathway, the ED pathway, the PP pathway, and the TCA cycle (Fig. 3). At two time points during exponential growth, we obtained similar metabolite 13C-labeling patterns, which confirmed pseudo-steady-state isotopic enrichment (Fig. 3). To elucidate the fluxes through 48 reactions in the metabolic pathway, we combined the metabolite labeling data with substrate consumption rates (accounting for the excretion of gluconate and 2-ketogluconate) and biomass growth rates (Fig. 4A; see also Fig. S2 and Tables S3 and S4). Adjusting for the carbon loss through metabolite secretions, approximately 64% of the glucose consumed from the extracellular medium was available for intracellular catabolism in *P. protegens* Pf-5. In conjunction with meeting the carbon mass balance constraints, the very close alignment between the MFA-estimated labeling patterns and those determined experimentally reflected the good quality of the optimization procedure for the model cellular fluxes (see Fig. S3).

(i) Involvement of glucose oxidation and the ED pathway. Glucose catabolism can be initiated via three different routes: direct phosphorylation to G6P, oxidation to gluconate in the periplasm before phosphorylation to 6P-gluconate, or additional periplasmic oxidation of gluconate to 2-ketogluconate before phosphorylation to 6P-gluconate (Fig. 1). Doubly 13C-labeled 6P-gluconate (86% to 89%) and doubly 13C-labeled G6P (63%) were both consistent with the assimilation of [1,2-13C2]glucose (Fig. 3A). In accordance with the ED pathway, wherein the first three carbons of 6P-gluconate become the doubly 13C-labeled pyruvate and the last three nonlabeled carbons become GAP, pyruvate was 50% to 53% doubly 13C-labeled, and DHAP (an isomer of GAP) was >95% nonlabeled (Fig. 3A). The 37% to 38% nonlabeled fraction of pyruvate was indicative of the nonlabeled fractions of precursor metabolites downstream of GAP (Fig. 3A). In the absence of the 6-phosphofructokinase enzyme in *P. protegens* Pf-5, and thus the lack of the traditional forward EMP pathway (7), the nonlabeled GAP and DHAP combined to produce the highly nonlabeled FBP (85% to 87%), which then cycled backward through the upper EMP pathway to result in the 67% to 70% nonlabeled F6P and 31% nonlabeled G6P (Fig. 3A). The MFA quantified the relative contribution of the three assimilation routes for the glucose-derived carbons in *P. protegens* Pf-5 (Fig. 4A). Only approximately 5% of the glucose uptake was directly converted to G6P in the cytosol, whereas up to 95% of glucose was oxidized to gluconate accompanied by another flux (12.7%) to 2-ketogluconate (Fig. 4A). Following the phosphorylation of the oxidized products of glucose to 6P-gluconate, our MFA determined that the flux through the ED pathway was 67.2% of the glucose uptake rate in *P. protegens* Pf-5 (Fig. 4A).

(ii) Oxidative versus non-oxidative PP pathway. Following a decarboxylation reaction through the oxidative PP pathway, doubly 13C-labeled 6P-gluconate would become singly 13C-labeled Ru5P, which would introduce singly 13C-labeled fractions into metabolites in the PP pathway (Fig. 3B). Thus, singly 13C-labeled fractions of xylulose 5-phosphate (Xu5P) (21% to 25%), ribose 5-phosphate (R5P) (22% to 25%), and sedoheptulose 7-phosphate (S7P) (20% to 23%) were due to flux through the oxidative PP pathway (Fig. 3B). However, by involving nonlabeled metabolites from downstream of the ED pathway, the non-oxidative PP pathway introduced relatively higher fractions of nonlabeled Xu5P (64% to 67%), R5P (65% to 69%), and S7P (62% to 69%) (Fig. 3B). Accordingly, the MFA determined an oxidative flux to the PP pathway (approximately 0.5%) that was a tenth of the non-oxidative PP pathway fluxes (5.1%) from ketolase and transaldolase reactions (Fig. 3B and 4A). As a precursor to both the oxidative PP
FIG 3 Long-term isotopic enrichment with [1,2-13C2]glucose for carbon mapping of the metabolic network structure in *P. protegens* Pf-5. Carbon mapping is illustrated on the left and the metabolite labeling data are provided on the right for the following: (A) Initial glucose catabolism, Embden-Meyerhof-Parnas (EMP) pathway, and the Entner-Doudoroff (ED) pathway; (B) oxidative and non-oxidative routes of the pentose-phosphate (PP) pathway; (C) the tricarboxylic acid (TCA) cycle. The dashed arrows describe minor formation routes of the metabolites. In the carbon mapping on the left, 13C carbons directly from glucose are in black, nonlabeled carbons from the ED pathway are in blue, singly 13C-labeled metabolites originated from the oxidative PP pathway are in orange, and the addition of labeled CO2 is shown by purple circles. Metabolite labeling patterns in panels A to C: nonlabeled (light blue), singly labeled (orange), doubly labeled (cream), triply labeled (green), and quadruply labeled (red). Labeling data (average ± standard deviation) were from independent biological replicates (*n* = 3). G6P, glucose 6-phosphate; 6P-Glucn, 6-phosphogluconate; F6P, fructose 6-phosphate; FBP, fructose 1,6-bisphosphate; DHAP, dihydroxyacetone-3-phosphate; GAP, glyceraldehyde 3-phosphate; PEP, phosphoenolpyruvate; 3-PG, 3-phosphoglycerate; Xu5P, xylulose 5-phosphate; R5P, ribose 5-phosphate; S7P, sedoheptulose 7-phosphate; OAA, oxaloacetate; Asp, aspartate; α-KG, α-ketoglutarate.
pathway and the ED pathway, 6P-gluconate represents an important branch-point metabolite in the network. Therefore, the low contribution of 6P-gluconate to the oxidative PP pathway necessitated a cyclic flux from the downstream ED pathway to the upper EMP pathway to channel glucose-derived carbons toward the non-oxidative PP pathway to support biomass growth (Fig. 4A).

(iii) Downstream metabolic pathways. The $^{13}$C-labeling patterns of TCA cycle intermediates were consistent with the established route of carbon flow through this pathway (Fig. 3C). The decarboxylation of pyruvate generated nonlabeled and singly $^{13}$C-labeled acetyl moieties in acetyl coenzyme A (acetyl-CoA), which were subsequently incorporated into the TCA cycle by combining with oxaloacetate (OAA; nonlabeled, singly, doubly, and minorly triply $^{13}$C labeled) to produce citrate (nonlabeled, singly, doubly, triply, and minorly quadruply $^{13}$C labeled) (Fig. 3C). The two sequential decarboxylation reactions in the TCA cycle led to the disappearance of the quadruply $^{13}$C-labeled fraction in citrate and, thereafter, the triply $^{13}$C-labeled fraction in α-ketoglutarate (Fig. 3C). The resulting succinate labeling pattern (nonlabeled, singly, and doubly $^{13}$C labeled) led to a similar labeling scheme through fumarate, malate, and

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**FIG 4** Quantitative metabolic flux analysis (A) and energetic distributions (B) in glucose-grown *P. protegens* Pf-5. All fluxes were normalized to glucose uptake, and the thickness of each arrow is scaled to the relative flux percentage. For panel A, black dotted arrows indicate contribution to biomass and gray dashed arrows indicate metabolite secretions. The absolute fluxes (average ± standard deviation), optimized on data obtained from independent biological replicates ($n = 3$), are listed in Table S3 and S4. Abbreviations in panel A are as listed in the legend of Fig. 3. For panel B, the relative contributions of different pathways to the NAD(P)H, UQH$_2$, and ATP pools were calculated from metabolic fluxes. Data for *P. putida* KT2440 were obtained from the metabolic flux analysis reported in Nikel et al. (10). All values in panel B are relative to glucose consumption rate and cellular biomass.
OAA (Fig. 3C). The MFA obtained a substantial flux (>70% of the glucose uptake) through the TCA cycle from OAA around to malate (Fig. 4A).

Anaplerotic reactions contributed to the triply $^{13}$C-labeled OAA (Fig. 3C). The aforementioned decarboxylation reactions in the TCA cycle contributed to the $^{13}$C-labeled carbon dioxide (CO$_2$) pool, which was calculated to be ~40% of the total dissolved CO$_2$ (Fig. 3; see also Fig. S4). The carboxylation of doubly $^{13}$C-labeled pyruvate or doubly $^{13}$C-labeled PEP with $^{12}$C-labeled CO$_2$ would generate triply $^{13}$C-labeled OAA (Fig. 3C). Notably, singly $^{13}$C-labeled OAA can be formed from carboxylation reactions of nonlabeled pyruvate or PEP with singly $^{12}$C-labeled CO$_2$, and from the generation of singly $^{13}$C-labeled malate though the TCA cycle using malate dehydrogenase (Fig. 3C). The relative contributions of the two precursors to OAA (i.e., pyruvate/PEP versus malate) in \textit{P. protegens} Pf-5 was resolved with the MFA, which determined a substantially higher fractional flux of pyruvate to OAA (70%) than the flux of malate to OAA (7.7%) (Fig. 4A). This low flux through malate dehydrogenase was accompanied by a high flux for the direct conversion of malate to pyruvate (63.4%), thus highlighting a very active pyruvate shunt in \textit{P. protegens} Pf-5 (Fig. 4A). Moreover, the $^{13}$C-labeling patterns of TCA cycle metabolites implied an inactive glyoxylate shunt, which bypasses the decarboxylation reactions in the canonical TCA cycle to produce malate and succinate from citrate (see Fig. S5). Specifically, there was a lack of triply $^{13}$C-labeled succinate, which would be produced from triply and quadruply $^{13}$C-labeled citrate through the glyoxylate shunt (Fig. S5).

(iv) Energetics of glucose catabolism. To capture possible species-dependent differences in energetics generated directly from the central carbon metabolism, we compared the yields of reduced ubiquinone (UQH$_2$), NAD(P)H, and ATP using our MFA-based cellular fluxes in \textit{P. protegens} Pf-5 and those previously reported for \textit{P. putida} KT2440, a well-studied biocatalyst candidate (10) (Fig. 4B). Compared to \textit{P. putida} KT2440 (10), there was a slightly higher flux (approximately 5% higher) from glucose to gluconate in \textit{P. protegens} Pf-5, but the flux from malate to OAA was lower (by approximately 25%) in \textit{P. protegens} Pf-5. Accordingly, there was a higher yield of UQH$_2$ from initial glucose catabolism in \textit{P. protegens} Pf-5 but a higher UQH$_2$ yield from the TCA cycle in \textit{P. putida} KT2440 (Fig. 4B). For the yield of NAD(P)H, there was a higher contribution from the oxidative PP pathway (by approximately 60%) in \textit{P. putida} KT2440 than in \textit{P. protegens} Pf-5, but the contribution of the flux from malate to pyruvate via the malic enzyme was lower (by approximately 81%) in \textit{P. putida} KT2440 than in \textit{P. protegens} Pf-5; the contribution of the TCA cycle to NAD(P)H generation remained about the same in both species (Fig. 4B). With respect to ATP production by substrate-level phosphorylation, \textit{P. protegens} Pf-5 produced less [by approximately 3 mmol ATP per cell dry weight in grams (g$_{CDW}$)] than \textit{P. putida} KT2440 because of the relatively lower fluxes in the downstream ED pathway of \textit{P. protegens} (10) (Fig. 4). However, the net ATP yields were similar, because \textit{P. protegens} Pf-5 consumed less ATP than \textit{P. putida} KT2440 in the initial glucose catabolism due to the higher flux of glucose oxidation to gluconate and 2-ketogluconate and the subsequent carbon loss due to the secretions of these oxidized products in \textit{P. protegens} Pf-5 (10) (Fig. 4). Remarkably, despite the different contributions of the relevant metabolic pathways, the combination of these contributions led to near-equivalent net yields of UQH$_2$, NAD(P)H, and ATP in \textit{P. protegens} Pf-5 and \textit{P. putida} KT2440 (Fig. 4B).

Hierarchy of glucose metabolism in the presence of other carbohydrates. (i) Proof of concept with $^{13}$C-labeled glucose and unlabeled glucose. Before determining the relative incorporation of \textit{[U-$^{13}$C$_6$]}glucose in the presence of a nonlabeled carbohydrate (xylose, arabinose, galactose, fructose, or mannose), we first conducted proof-of-concept experiments with the cells grown on \textit{[U-$^{13}$C$_6$]}glucose alone or in a 1:1 mixture with unlabeled glucose (Fig. 5 and 6) (24). By a comparative analysis with the metabolite labeling patterns in the latter two conditions, we sought to determine the relative incorporation of other unlabeled carbohydrates in the presence of \textit{[U-$^{13}$C$_6$]}glucose (Fig. 5 and 6).
Glucose with xylose, arabinose, or galactose. During growth on $^{13}$C-labeled glucose and galactose, the labeling patterns of metabolites in the EMP and ED pathways (F6P, FBP, and DHAP) were identical to the metabolite labeling during feeding on glucose alone, thus indicating the lack of galactose catabolism in the presence of glucose (Fig. 5A). During simultaneous feeding on $^{13}$C-labeled glucose and a pentose substrate (xylose or arabinose), the labeling patterns of metabolites in the PP pathway (Xu5P, R5P, and S7P) also indicated the lack of pentose assimilation (Fig. 5B). Furthermore, we found no incorporation of unlabeled xylose into $\alpha$-ketoglutarate, consistent with the lack of the Weimberg pathway (see Fig. S6). The previously reported absence of $P.\ protegens$ Pf-5 growth on medium containing only galactose, xylose, or arabinose implied the absence of assimilation of these carbohydrates (6). However, a recent study (21) showed extracellular xylose depletion by $P.\ protegens$ Pf-5 during growth on a mixture of carbohydrates, suggesting that a lack of growth is not predictive of extracellular substrate depletion or possible assimilation. Our $^{13}$C-labeling data explicitly ascertained the absence of carbon assimilation from galactose, xylose, or arabinose in the presence of glucose (Fig. 5).

![FIG 5 Metabolite labeling patterns during growth on [U-$^{13}$C$_6$]glucose ($^{13}$C$_6$-Glucose) alone or with unlabeled glucose ($^{12}$C$_6$-Glucose), unlabeled galactose ($^{12}$C$_6$-Galactose), unlabeled xylose ($^{12}$C$_6$-Xylose), or unlabeled arabinose ($^{12}$C$_6$-Arabinose). (A) Carbon mapping (left) and the labeling data (right) for intracellular metabolites in the upper Embden-Meyerhof-Parnas pathway following feeding on $^{[13C_6]}$glucose alone or with $^{[12C_6]}$glucose or $^{[2C_6]}$galactose. (B) Carbon mapping (left) and the labeling data (right) for intracellular metabolites in the pentose-phosphate pathway following feeding on $^{[13C_6]}$glucose alone or with $^{[12C_6]}$glucose, $^{[12C_6]}$xylose, or $^{[12C_6]}$arabinose. In the carbon mapping in panels A and B, the open circles and the filled circles represent nonlabeled and $^{13}$C-labeled carbons, respectively. Data were obtained at two time points during exponential growth: at OD$_{600}$ 0.5 to 0.6 (T1) and at OD$_{600}$ 0.9 to 1.0 (T2). Metabolite labeling data (average ± standard deviation) were from independent biological replicates ($n = 3$). Very small error bars are not noticeable. The metabolite abbreviations are as listed in legends for Fig. 1 and 3.]
(iii) Glucose with fructose. Following growth simultaneously on \([U-^{13}C_6]glucose\) and unlabeled fructose, there was a persistent presence of nonlabeled fractions in the intracellular metabolites, indicating both uptake and assimilation of fructose in the presence of glucose (Fig. 6). However, the high abundance of the nonlabeled fraction and the increased pool size of FBP compared to that in cells grown only on glucose indicated a bottleneck in fructose assimilation, which may explain the lower rate of fructose uptake (2.02 ± 0.86 mmol g_{CDW}^{-1} \cdot h^{-1}) than glucose uptake (7.01 ± 1.70 mmol g_{CDW}^{-1} \cdot h^{-1}) (Fig. 2B and 6; see also Fig. S7). In accordance with fructose incorporation through F1P into FBP, the highest fraction of nonlabeled carbons

FIG 6 Metabolite labeling patterns during growth on \([U-^{13}C_6]glucose\) alone or with unlabeled glucose \((^{12}C_6-Glucose)\), unlabeled fructose \((^{12}C_6-Fructose)\), or unlabeled mannose \((^{12}C_6-Mannose)\). Carbon mapping (left) and the labeling data (right) for intracellular metabolites in the upper Embden-Meyerhof-Parnas (EMP) pathway, the pentose-phosphate pathway, and the Entner-Doudoroff pathway. The open circles and the filled circles represent nonlabeled and \(^{13}C\)-labeled carbons, respectively; the green circles represent labeling schemes specifically from reverse flux through upper EMP pathway. Data were obtained at two time points during exponential growth: at OD_{600} of 0.5 to 0.6 (T1) and at OD_{600} of 0.9 to 1.0 (T2). NF, not found. Labeling data (average ± standard deviation) were from independent biological replicates (n = 3). Very small error bars are not noticeable. The abbreviations are as listed in the legends for Fig. 1 and 3.
was in FBP (59% to 62%); both F6P and DHAP had lower nonlabeled fractions (25% to 30% and 17%, respectively) (Fig. 6). The lower fraction of nonlabeled carbons in DHAP than in F6P implied that the fructose-derived carbons were preferentially routed via a backward flux through upper EMP pathway (i.e., from FBP to G6P) toward the ED pathway (Fig. 6). The labeling of G6P reflected the nonlabeled and partially 13C-labeled fractions from F6P, consistent with this backward flux (Fig. 6). Therefore, in lieu of the forward EMP pathway (i.e., from G6P to FBP), our data stressed the importance of the ED pathway in the co-processing of fructose with glucose, and this was further verified below with 13C-MFA.

The labeling patterns of metabolites in the PP pathway also showed that the contribution of the fructose-derived carbons in this pathway was preferentially through the non-oxidative route, as previously shown in cells grown on glucose alone (Fig. 3 and 6). A transketolase reaction in the non-oxidative PP pathway combines the first two carbons of F6P with GAP to produce Xu5P. Both Xu5P and R5P have significant fractions of triply 13C-labeled carbons (38% to 44%), in accordance with the combination of nonlabeled F6P with triply 13C-labeled GAP following growth on 13C-labeled glucose with unlabeled fructose (Fig. 6). Due to the low fraction of nonlabeled GAP (as determined from DHAP labeling), there was a lack of doubly 13C-labeled R5P and Xu5P, which was also evident in cells grown on [U-13C6]glucose with unlabeled fructose (Fig. 6).

(iv) Glucose with mannose. In contrast to the metabolite labeling during growth on [U-13C6]glucose with unlabeled fructose, the metabolite labeling patterns following feeding on the [U-13C6]glucose with unlabeled mannose showed delayed assimilation of mannose into intracellular metabolism during exponential cell growth (Fig. 6). This was evident by the substantial increase in the nonlabeled fraction of FBP across the two time points, from 12% at an optical density at 600 nm (OD600) of ~0.5 to 52% at OD600 of ~1.0 (Fig. 6). This time-dependent labeling data matched with the initiation of a significant decrease in extracellular mannose concentration after 6 h of growth following the depletion of glucose (Fig. 2B).

A higher nonlabeled fraction of FBP than of F6P implied that the incorporation of mannose-derived carbons into FBP by way of fructose was preferred in P. protegens Pf-5 (Fig. 6). In agreement with this catabolic route for mannose, there was a larger pool of FBP than F6P when cells were grown on a mixture of glucose and mannose relative to when cells were grown on glucose only—this relatively larger FBP pool was also obtained in cells grown on a mixture of glucose and fructose (Fig. S7). Following growth on [13C]glucose with unlabeled mannose, the nonlabeled fraction of DHAP (18%) was lower than the corresponding fraction in F6P (28%) and G6P (26%) (Fig. 6). These labeling data indicated collectively a cycling of nonlabeled mannose carbons backward through the upper EMP pathway toward the ED pathway, similar to the intracellular metabolic route of fructose (Fig. 6).

Cellular carbon fluxes during co-utilization of glucose and fructose. Quantitative MFA was used to assess the differences in the metabolic fluxes during growth on glucose alone versus growth on glucose with fructose (Fig. 7; see also Tables S4 and S5). Our MFA focused on the pathways surrounding the cellular entry points of carbohydrates: initial glucose catabolism, the EMP pathway, the PP pathway, and the ED pathway (Fig. 7). Each MFA made use of the labeling data collected at an OD600 of 0.5 during early exponential phase before extracellular glucose is depleted and was constrained by the consumption rate of each carbohydrate, biomass effluxes, and metabolite secretions (Fig. 7 and S2; Tables S5 and S6). Upon optimization of the estimated metabolic fluxes, the model-estimated 13C-labeling patterns agreed well with the experimentally determined 13C-labeling patterns for each condition (see Fig. S8 and S9).

Despite the genome-encoded capabilities in P. protegens Pf-5 to split FBP into DHAP and GAP (7), the MFA revealed that the net flux was instead the aldolase reaction that combines DHAP and GAP to generate FBP (Fig. 7). Due to the additional incorporation
of fructose-derived carbons through the flux from FBP to 6P-gluconate, there were higher fluxes from FBP to F6P (3-fold increase), F6P to G6P (3.6-fold increase), and G6P to 6P-gluconate (2.3-fold increase) in cells grown on the mixture of glucose and fructose compared to those grown on glucose alone (Fig. 7). Similar to the metabolism of glucose alone, there was a significant flux of the glucose uptake channeled through glucose oxidation to gluconate (88%) and the ED pathway (95%) during the metabolism of both glucose and fructose (Fig. 7). Consistent with the increased carbon flux toward 6P-gluconate from fructose assimilation, there was a 4.3-fold increase in the flux toward the oxidative PP pathway (i.e., from 6P-gluconate to Ru5P) and a 19% increase in the ED pathway (i.e., from 6-gluconate to GAP and pyruvate) (Fig. 7).

**Kinetic isotope analysis during co-utilization of glucose and mannose.** Due to the lack of isotopic pseudo-steady state in metabolite labeling during growth on the glucose-mannose mixture, 13C-MFA could not be performed. To capture the assimilation route for mannose, we obtained labeling measurements at six time points during exponential growth on 13C-labeled glucose and unlabeled mannose (Fig. 8A; see also Fig. S10). Specifically, we examined the labeling patterns of gluconate, 6P-gluconate, G6P, F6P, FBP, and DHAP (Fig. 8A and S10). Across all time points, gluconate labeling was consistently ~100% fully 13C labeled, indicating that gluconate was made exclusively from the 13C-labeled glucose (Fig. S10). However, the other five metabolites had
Amongst all the metabolites, 6P-gluconate exhibited the slowest kinetic incorporation of nonlabeled fractions (Fig. S10). The appearance of a nonlabeled fraction in 6P-gluconate (starting at 4%) occurred at an OD$_{600}$ of 0.94, the fifth time point (Fig. S10). In contrast, FBP exhibited the fastest incorporation of a nonlabeled fraction starting at an OD$_{600}$ of 0.2 (Fig. 8A and S10). Compared to the labeling kinetics of FBP, there was a delay in the incorporation of nonlabeled carbons in F6P and G6P, which started to occur at an OD$_{600}$ of 0.6, and DHAP, which steadily increased after an OD$_{600}$ of 0.8 (Fig. 8A and S10). To determine whether there was statistical significance regarding how mannose-derived nonlabeled carbons were incorporated into metabolism, we analyzed the rates at which nonlabeled carbons were incorporated into FBP, F6P, G6P, and DHAP with a mixed-effects model. This statistical analysis ($F_{3,57} = 12.973$, $P < 0.0001$) confirmed that the significant effect of OD$_{600}$ on the incorporation of mannose-derived nonlabeled carbons was dependent on the metabolite (Fig. 8A). Collectively, these kinetics data demonstrated that, instead of being channeled directly from FBP to GAP and DHAP, mannose carbons were incorporated at FBP and cycled up through F6P and G6P toward the ED pathway to generate subsequently GAP and DHAP (Fig. 8). Thus, the catabolic route for mannose was similar to what was determined for fructose catabolism.

**FIG 8** Metabolic routing of mannose through central metabolism in the presence of glucose. (A) Kinetic profiling of metabolite labeling patterns in cells grown simultaneously on [U-13C$_6$]glucose and unlabeled mannose; the lines through the data are to guide the eye. (B) Dynamic flux ratio analysis for the metabolic entry point of glucose versus mannose assimilation through G6P and FBP, respectively: flux from mannose (light orange), flux from ED pathway (gray), flux from F6P (green), flux from glucose (blue). Data in panel B were obtained at six time points during exponential growth: at OD$_{600}$ of 0.21 to 0.27 (T1), at OD$_{600}$ of 0.40 to 0.44 (T2), at OD$_{600}$ of 0.54 to 0.67 (T3), at OD$_{600}$ of 0.79 to 0.87 (T4), at OD$_{600}$ of 0.90 to 0.98 (T5), and at OD$_{600}$ of 1.2 to 1.4 (T6). In panels A and B, The data (average ± standard deviation) are from biological replicates ($n = 3$). The metabolite abbreviations are as listed in the legend for Fig. 1.
Using metabolic flux ratio analysis across the different time points, we quantified the relative incorporation of glucose versus mannose at relevant entry points (G6P and FBP, respectively) into the metabolic network (Fig. 8B). The metabolite G6P is a result of fluxes from both direct glucose phosphorylation and from F6P derived from the cyclic ED-EMP pathway flux; the metabolite FBP is synthesized either from mannose or from downstream ED pathway flux through the combination of GAP and DHAP (Fig. 8B). In agreement with the extracellular depletion profiles (Fig. 2B), the fractional flux of glucose directly to G6P started at 81% at an OD$_{600}$ of 0.2 and steadily decreased over time to reach eventually 2% at an OD$_{600}$ of 1.2 (Fig. 8B). Conversely, the relative contribution of mannose-derived FBP went from 5% to eventually 64% at the highest measured OD$_{600}$ (Fig. 8B). The cycling of carbons from mannose toward the ED pathway maintained a 36% fractional flux of triose phosphates to FBP, even after glucose was depleted (Fig. 8). In sum, the metabolic flux ratio analysis demonstrated simultaneous assimilation of both glucose and mannose into the intracellular metabolites, but the relative contribution of each substrate varied as a function of time (Fig. 8B).

**DISCUSSION**

The metabolic networks and fluxes of several *Pseudomonas* species, including *P. putida*, *P. fluorescens*, and *P. aeruginosa*, have been previously described (8–11, 22, 25). Here, we present the first metabolic flux analysis of the recently characterized *P. protegens* PF-5, a common plant commensal bacterium known to secrete specialized metabolites important for the biocontrol of fungi and bacteria pathogenic to plants (4, 6). Metabolic flux quantitation determined that initial glucose catabolism in *P. protegens* PF-5 was primarily through periplasmic oxidation to gluconate, with relatively minor influx of glucose through G6P (Fig. 4A). Up to 95% of consumed glucose in *P. protegens* PF-5 was channeled through the ED pathway, which was also reported in *P. putida*, *P. fluorescens*, and *P. aeruginosa* (8, 11, 22, 25). Further glucose catabolism was routed through a cyclic flux from the ED pathway to the upper EMP pathway, which was also elucidated in *P. putida* KT2440 (10, 11). Furthermore, the non-oxidative route was more significant than the oxidative route in generating PP pathway intermediates in *P. protegens* PF-5, as previously reported for *P. putida* KT2440 (11) (Fig. 3B and 4A). The highly active flux of pyruvate formation from malate in the *P. protegens* PF-5 cells was also reported in *P. putida* and *P. fluorescens* (8, 10, 22). In sum, our results stressed that the metabolic network for glucose metabolism in *P. protegens* PF-5 is consistent with the metabolic network of previously studied species of the *Pseudomonas* genus. Finally, through the contribution of different metabolic pathways, the total UQH$_2$, NAD(P)H, and ATP yields of *P. protegens* PF-5 were remarkably similar to those of *P. putida* KT2440, whose metabolism has been featured for its capability of fulfilling high demands of reducing power (10).

Root exudates and the breakdown of polysaccharides from plant biomass both provide various carbohydrates that stimulate the growth of soil microorganisms, including *Pseudomonas* species (26). With respect to the catabolism of other carbohydrates besides glucose, we found that *P. protegens* PF-5 did not metabolize the common hemicellulose monomers galactose, xylose, or arabinose in the presence of glucose but did utilize the carbon mixtures of glucose with fructose or mannose (Fig. 5 and 6). The lack of assimilation of the pentoses and galactose was consistent with the absence of genes for the relevant catabolic pathways. The preference for glucose over fructose and mannose implied catabolite repression by glucose. An elucidation of the underlying mechanism of this repression requires further transcriptional expression and mutation studies. In accordance with gene annotations, metabolite labeling data confirmed that fructose was incorporated into metabolism via FBP (Fig. 6). However, contrary to the possible route for mannose assimilation through F6P annotated in the *P. protegens* PF-5 genome (7), the primary route of mannose assimilation in *P. protegens* PF-5 was found to be also via FBP (Fig. 6). In addition, the appearance of fructose extracellularly during growth on the mixture of glucose and mannose implied a conversion of mannose to...
fructose prior to intracellular metabolism (Fig. 2B). Mannose conversion to fructose by a mannose isomerase has been reported previously in *P. cepacia*, *P. aeruginosa*, and *P. saccharophila* (17–20). Whether a nonspecific isomerase exists in *P. protegens* Pf-5 remains to be determined.

The cyclic metabolism linking backward flux from FBP through the upper EMP pathway toward the ED pathway produces equivalent amounts of NADPH, but less net ATP, compared to that with direct FBP conversion to GAP and DHAP. Interestingly, instead of this direct contribution to the lower EMP pathway, carbons from assimilated fructose and mannose were routed through the cyclic connection between the upper EMP pathway and the ED pathway during mixed-hexose utilization (Fig. 7 and 8). Comparable MFA findings were reported for fructose-only catabolism in *P. putida* KT2440 and *P. fluorescens* SBW25 (14, 16). Therefore, *P. protegens* Pf-5 exhibits a strong reliance on the ED pathway for both fructose and mannose assimilation even in the presence of glucose.

Similar growth phenotypes during co-utilization of hexoses implied that, despite different carbohydrates in the growth medium at the same total carbon content, *P. protegens* Pf-5 preserved a constant biomass (Fig. 2A). However, the uptake of glucose was preferred over the uptake of fructose (3 to 1) or mannose (4 to 1) (Fig. 2; see also Table S2 in the supplemental material). The composition ratio of glucose to fructose in maize root exudates was found to be 2 to 1 (27). Across soil horizons, the glucose-mannose ratio ranged approximately from 3:1 to 5:1 (28). Therefore, our relative consumption rates of glucose versus fructose or mannose in *P. protegens* Pf-5 were remarkably in agreement with the relative composition of these carbohydrates in environmentally relevant conditions. Whether this agreement is due to an evolutionary connection to nutritional adaptability warrants further investigation.

In regards to the potential of *P. protegens* Pf-5 as a biocatalytic platform, the innate production and secretion of gluconate and 2-ketogluconate in *P. protegens* Pf-5 are attractive features. Oxidized sugars are important precursors to polymeric materials, including polyesters. In fact, gluconate was identified as a top 30 value-added candidate for the production of bio-inspired materials (29). Under our experimental conditions, the quantitative flux analysis determined that the secretion rates of gluconate and 2-ketogluconate collectively accounted for approximately 35% of the glucose uptake in *P. protegens* during exponential growth, whereas metabolite secretion of gluconate was reported to be less than 10% of the glucose uptake in *P. putida* KT2440 (11) (Fig. 2C). While there were relatively similar secretions during growth on glucose alone or on glucose and fructose, growth on glucose and mannose resulted in a decrease in the total secretion (by approximately 3.5 mM) of gluconate and 2-ketogluconate (Fig. 2C). After glucose was depleted, decreases in the concentrations of both gluconate and 2-ketogluconate indicated that the cells can utilize these metabolites once their favored carbon source (i.e., glucose) was absent (Fig. 2B and C). This phenomenon would need to be considered and manipulated to harvest these metabolite secretions in *P. protegens* Pf-5 as valuable products. In light of the abundance of different types of carbohydrates in natural environments and in renewable carbon feedstocks used for engineered bioproduction (30), our findings collectively provide important insights regarding the cellular metabolism underlying carbohydrate co-utilization in *P. protegens* Pf-5 and related *Pseudomonas* species.

**MATERIALS AND METHODS**

**Materials.** The *P. protegens* Pf-5 cells were acquired from the American Type Culture Collection (Manassas, VA). Unless noted otherwise, the chemicals used in the growth media were obtained from Sigma-Aldrich (St. Louis, MO), Cayman Chemical (Ann Arbor, MI), or Fisher Scientific (Pittsburgh, PA). The 13C-labeled glucose ([U-13C6]glucose and [1,2-13C6]glucose) were purchased from Cambridge Isotopes (Tewksbury, MA) and Omicron Biochemicals (South Bend, IN), respectively. All culture solutions were prepared with Millipore water (18.2 MΩ·cm; Millipore, Billerica, MA, USA) while resuspensions for LC-HRMS analysis were made with LC-MS-grade water. Solutions were sterilized by passing through a 0.22-µm nylon filters (Waters Corporation, MA). An Agilent Cary UV-visible spectrophotometer (Santa Clara, CA) was used for optical density readings at 600 nm. The LC-HRMS analysis was conducted on an ultra-high-performance LC (Dionex UltiMate 3000; Thermo Scientific) coupled to a high-resolution
Culturing conditions and growth measurements. Batch growth experiments (three to seven biological replicates) of _P. protegens_ PF-5 were conducted in an incubator (model I24; New Brunswick Scientific, Edison, NJ) maintained at 30°C and shaken at 220 rpm. The initial growth in nutrient-rich medium prior to growth in minimal-nutrient medium was conducted as previously described (11). Final growth experiments were conducted in 125-ml baffled flasks with a pH-adjusted (7.0) filter-sterilized minimal-nutrient medium that contained major salts and essential trace metal nutrients as previously reported (31): 89.4 mM K_HPO_4, 56.4 mM NaH_2PO_4, 0.81 mM MgSO_4·7H_2O, 18.7 mM NH_4Cl, 8.6 mM NaCl, 34 μM CaCl_2·2H_2O, 30 μM FeSO_4·7H_2O, 0.86 μM CuSO_4·5H_2O, 1.9 μM HBO_3, 7.7 μM ZnSO_4·7H_2O, 0.75 μM MnSO_4·H_2O, 0.26 μM NiCl_2·6H_2O, and 0.3 M Na_2MoO_4·5H_2O. The carbohydrate composition was 100 mM C total for glucose alone (equivalent to 16.7 mM or 3 g·liter⁻¹ glucose) and for 1:1 mixtures of glucose and xylose, glucose and arabinose, glucose and galactose, glucose and mannose, or glucose and fructose. For cellular isotopic enrichment, either [U-¹³C₆]glucose or [1,2-¹³C₆]glucose was used in glucose-only growth, but only [U-¹³C₆]glucose was used for mixtures in combination with an unlabeled second carbohydrate. Bacterial growth in the biological replicates was monitored as a function of time until late stationary phase using OD₆₀₀ measurements (see Fig. S1 in the supplemental material)—cell suspensions were diluted when the OD₆₀₀ value was above 0.5 to get accurate readings. Cell dry weight in grams was also determined throughout growth by lyophilizing the cell pellets as previously described (11). Biomass growth rate (per hour) for each growth condition was obtained by regression analysis of OD₆₀₀ measurements over time (Table S2).

Measurement of carbohydrate consumption. Independent ¹³C-tracer experiments, as described in “Metabolic monitoring and quantification,” confirmed that extracellular depletion of the substrates correlated with metabolic activity. The extracellular depletion of each carbohydrate substrate (three biological replicates) was determined throughout 24 h of cell growth. Culture aliquots were pelleted by centrifugation, and the supernatants were stored at −20°C until further analysis. According to previously reported LC methods for carbohydrate analysis (21), we applied an analytical method using LC-HRMS for monitoring the carbohydrate concentration in the extracellular solution. Peak identification and quantification of carbohydrate concentrations were conducted with Thermo Scientific Xcalibur 3.0 Quan Browser. Carbohydrate consumption rates (in millimoles per gram [dry weight] per hour) were obtained by combining the regression analyses of carbohydrate depletion over time with biomass growth rates (Table S2).

Metabolite monitoring and quantification. (i) Extracellular metabolites. To determine metabolite extrusion rates, cell suspensions were harvested periodically throughout growth and pelleted with centrifugation, and the supernatants were removed and stored at −20°C until LC-HRMS analysis. Dilutions of 1:10, 1:100, and 1:1,000 were conducted to account for the various concentrations of each metabolite over time. For the LC, an Acquity UPLC Waters 1.7-μm particle size column with dimensions of 2.1 mm by 100 mm was used for all metabolomics samples (Milford, MA), with a constant column temperature of 25°C. The flow rate was kept constant at 0.180 ml·min⁻¹. The mobile phase composition and LC protocol were as previously described (31). The injection volume for each sample was 10 μL. The MS was operated in full-scan negative mode. Metabolite identification was based on accurate mass and matches with standard retention times. Metabolite levels were quantified using Thermo Scientific Xcalibur 3.0 Quan Browser.

(ii) Intracellular metabolites. Cells were separated by filtration and then lysed to extract the intracellular metabolites as described in Sasnow et al. (11). Metabolites in solution were monitored by LC-HRMS, and the ¹³C labeling patterns were analyzed on Metabolomic Analysis and Visualization Engine (MAVEN) software (32, 33). Isotopologue data were obtained for the following compounds: 6P-glucose, G6P, F6P, FBP, DHAP, 3-phosphoglycerate, PEP, pyruvate, Xu5P, R5P, S7P, aspartate, citrate, α-ketoglutarate, succinate, and malate. Aspartate ¹³C labeling was used as a proxy for OAA ¹³C labeling by assuming equilibrium between the two compounds (11). The labeling of dissolved CO₂ was estimated from the labeling patterns of ornithine and citrulline (Fig. S2); ornithine incorporates one mole of dissolved CO₂ to become citrulline. All the extracted isotopologues were corrected for natural abundance of ¹³C. To verify pseudo-steady-state isotopic enrichment of the intracellular pools, metabolites were isolated from cellular extracts obtained at two different time points during the exponential phase, at OD₆₀₀ values of ~0.5 and ~1.0 (11). To analyze mannose incorporation over time, cells were extracted at six time points during exponential growth corresponding to OD₆₀₀ values of ~0.2, ~0.4, ~0.6, ~0.8, ~0.9, and ~1.3. A mixed-effects model was used for analyzing R (34) and the lmerTest package (35), which modeled the nonlabeled fraction (log transformed) by OD₆₀₀ metabolite, and their interaction with the random effect of biological replicates.

Quantitative metabolic flux modeling. The quantitation of the metabolic fluxes was achieved for cells grown on [1,2-¹³C₂]glucose alone, [U-¹³C₆]glucose and unlabeled glucose, or [U-¹³C₆]glucose and unlabeled fructose. Additionally, we employed the following experimental data to constrain the metabolic flux analysis: substrate uptake rates, metabolite excretions, growth rate, and cellular stoichiometry. To determine the uptake of glucose-derived carbons from the periplasm to the cytosol (to add as constraints in the flux modeling), a regression analysis was conducted on the extracellular glucose depletion after subtracting the extracellular amounts of both glucose and 2-ketogluconate (Table S2; Fig. 2). We constrained the model to account for periplasmic oxidation reactions and secretion rates based on the adjusted substrate uptake rates into cytosol (Fig. S2). Carbon effluxes from intermediates in central metabolism toward biomass production were determined on the basis of the growth rate under each condition and the biomass composition of _P. putida_ (nucleic acids, proteins, cell membrane,
and carbohydrate polymers) (36). An initial reaction network for the central carbon metabolism of _P. putida_ PF-5 was constructed using a predicted genome-scale metabolic model (7) and gene annotation of metabolic enzymes reported on the KEGG database (37–39) and MetaCyc (40). The metabolic reaction network was validated through $^{13}$C labeling of intracellular metabolites. The following reactions were constrained in the forward direction: gluconate $\rightarrow$ 6P-gluconate, 6P-gluconate $\rightarrow$ ribulose 5-phosphate, gluconate $\rightarrow$ 2-ketogluconate, 2-ketogluconate $\rightarrow$ 6P-gluconate, glucose $\rightarrow$ G6P, FBP $\rightarrow$ F6P, malate $\rightarrow$ pyruvate, pyruvate $\rightarrow$ OAA, and PEP $\rightarrow$ OAA. Optimized fluxes in the model metabolic network reactions were determined by the 13CFLUX2 software package (http://www.13cflux.net) (41), whereby the quality of fit was optimized iteratively by comparing experimental $^{13}$C-labeling data and the in silico-estimated labeling data.

For cells grown on $[U-^{13}$C$_6]$glucose and mannose, we conducted a metabolic flux ratio analysis across six time points during exponential growth (see Fig. S11). Specifically, we determined the relative flux contributions of substrate uptake versus intracellular metabolic reactions to the biosynthesis of G6P and FBP (Fig. S11). Optimized flux ratios were obtained by iterative optimization with the objective of obtaining the minimum error difference between experimental and simulated labeling data (Fig. S11).

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at https://doi.org/10.1128/AEM.02084-18.

**SUPPLEMENTAL FILE 1**, PDF file, 1 MB.

**ACKNOWLEDGMENTS**

Graduate financial support for R.A.W. and C.M.M. was provided by the College of Agriculture and Life Sciences at Cornell University.

We thank Krista A. Barzen-Hanson from the Aristilde Research Group at Cornell University (presently at Elmira College, Division of Mathematics and Natural Sciences) for her technical assistance in implementing the LC-MS protocol to analyze the carbohydrate compounds in solution. We acknowledge Erika Mudrak at the Cornell Statistical Consulting Unit for her assistance with implementing and interpreting the statistical analysis.

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