Nanopore Long-Read Guided Complete Genome Assembly of *Hydrogenophaga intermedia*, and Genomic Insights into 4-Aminobenzenesulfonate, *p*-Aminobenzoic Acid and Hydrogen Metabolism in the Genus *Hydrogenophaga*

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We improved upon the previously reported draft genome of *Hydrogenophaga intermedia* strain PBC, a 4-aminobenzenesulfonate-degrading bacterium, by supplementing the assembly with Nanopore long reads which enabled the reconstruction of the genome as a single contig. From the complete genome, major genes responsible for the catabolism of 4-aminobenzenesulfonate in strain PBC are clustered in two distinct genomic regions. Although the catabolic genes for 4-sulfocatechol, the deaminated product of 4-aminobenzenesulfonate, are only found in *H. intermedia*, the *sad* operon responsible for the first deamination step of 4-aminobenzenesulfonate is conserved in various *Hydrogenophaga* strains. The absence of *pabB* gene in the complete genome of *H. intermedia* PBC is consistent with its *p*-aminobenzoic acid (pABA) auxotrophy but surprisingly comparative genomics analysis of 14 *Hydrogenophaga* genomes indicate that pABA auxotrophy is not an uncommon feature among members of this genus. Of even more interest, several *Hydrogenophaga* strains do not possess the genomic potential for hydrogen oxidation, calling for a revision to the taxonomic description of *Hydrogenophaga* as “hydrogen eating bacteria.”

**Keywords:** *Hydrogenophaga*, Nanopore, comparative genomics, hydrogen metabolism, phylogeny

**INTRODUCTION**

The genus *Hydrogenophaga* consists of rod-shaped yellow-pigmented gram negative bacteria that are generally considered capable of using hydrogen as an energy source (Willems et al., 1989). Among all currently described species of *Hydrogenophaga*, *Hydrogenophaga intermedia* is one of the most studied species due to its distinctive ability to efficiently degrade 4-aminobenzenesulfonate (4-aminobenzenesulfonate), a recalcitrant intermediate compound in
the synthesis of colorants (Dangmann et al., 1996). The catabolism of 4-aminobenzenesulfonate in H. intermedia S1 is by far the most well-studied among all described 4-aminobenzenesulfonate degraders (Perèi et al., 2001; Wang et al., 2009; Gan et al., 2011b; Yang et al., 2012; Hayase et al., 2016). H. intermedia S1 was isolated from wastewater and could grow on 4-aminobenzenesulfonate as the sole carbon source in a two-species bacterial coculture with Agrobacterium radiobacter S2 (Dangmann et al., 1996). To date, several enzymes associated with the downstream degradation of 4-aminobenzenesulfonate in strain S1 have been identified, cloned, characterized and functionally validated (Contzen et al., 2001; Halak et al., 2006, 2007). However, the first deamination step converting 4-aminobenzenesulfonate to 4-sulfocatechol, which was presumed to be catalyzed by a dioxygenase is yet to be indentified in strain S1 (Contzen et al., 2001; Halak et al., 2006, 2007).

More than 20 years after the isolation of the S1/S2 mixed culture, a second 4-aminobenzenesulfonate-degrading two-species bacterial culture consisting of Hydrogenophaga sp. PBC and Rabstonia sp. PBA isolated from textile wastewater was reported (Gan et al., 2011b). Transposon mutagenesis of strain PBC led to the identification of the sad operon that is responsible for the initial deamination step of 4-aminobenzenesulfonate to 4-sulfocatechol. The deamination step was presumed to operate similarly H. intermedia strain S1 given its near identical 16S rRNA sequence to strain PBC (Gan et al., 2011a,b, 2012c). In addition, both Hydrogenophaga strains could only be grown in axenic culture with 4-aminobenzenesulfonate as the sole carbon and nitrogen source if they were supplemented with p-aminobenzoate and biotin (Kampfer et al., 2005; Gan et al., 2011a,b), providing insight into the syntrophic relationship between the Hydrogenophaga strains and their helper strains in addition to suggesting that the biosynthesis pathways for these compounds were either missing or non-functional in H. sp. PBC and H. intermedia S1 (Dangmann et al., 1996; Gan et al., 2011b).

Strain PBC was the first Hydrogenophaga strain to have its genome sequenced (Gan et al., 2012b). However, due to the short read length of Illumina sequencing (2 × 100 bp) at that time and the high genomic GC content of strain PBC, the assembled genome was relatively fragmented. Further, most of the genes coding for 4-aminobenzenesulfonate metabolism were located on different contigs, limiting the analysis of the genomic structure and regulation of this pathway (Hegedüs et al., 2017). Interestingly, genes coding for biosynthesis of biotin and 4-aminobenzoic acid could not be identified in the draft genome of strain PBC, which is consistent with its requirement for these supplementalations for growth in axenic culture (Gan et al., 2011b; Kim and Gan, 2017). However, given that gaps exist in the draft genome, it may be possible that these gene regions were not assembled, thus emphasizing the need for a complete genome assembly to verify this observation. As more Hydrogenophaga genomes became available, H. sp. PBC was recently reclassified as H. intermedia PBC based on in silico genome–genome hybridization (Kim and Gan, 2017). Interestingly, despite the availability of various whole genome sequences for the type species of Hydrogenophaga, comparative genomic analyses of Hydrogenophaga currently are limited and most analyses have focused on a single genome (Gan et al., 2011b; Kim and Gan, 2017).

One of the defining traits of the genus Hydrogenophaga is the ability to grow chemorganotrophically or chemolithoautotrophically by oxidizing hydrogen as an energy source (Willems et al., 1989). A [NiFe]-membrane-bound hydrogenase has been isolated and purified from H. sp. AH-42 providing the first biochemical characterization of hydrogenase and molecular insight into the genetic components responsible for hydrogen oxidation in the genus Hydrogenophaga (Yoon et al., 2008, 2009). However, exceptions to the defining feature of the genus, e.g., inability to oxidize hydrogen under standardized culture condition, have been reported in two Hydrogenophaga type strains namely H. intermedia S1T and H. atypica DSM 15342T (Contzen et al., 2000; Kampfer et al., 2005), naturally inviting a comprehensive genomic survey of hydrogen-oxidizing genes in all sequenced members of the genus Hydrogenophaga.

To improve the genome assembly of H. intermedia PBC, we used the Nanopore MinION portable long-read sequencer to generate long sequencing reads coupled with a hybrid assembly using the Illumina dataset generated previously by Gan et al. (2012b). We showed that the incorporation of the Nanopore long reads enabled the successful assembly of H. intermedia PBC genome into a single contig. Further, we also provided an updated whole genome-based phylogeny of the family Comamonadaceae in addition to performing comparative genomics analysis of the genus Hydrogenophaga for the first time focusing on the prevalence of genomic potential for hydrogen oxidation, pABA synthesis and 4-aminobenzenesulfonate biodegradation.

**MATERIALS AND METHODS**

**Illumina and Nanopore Whole Genome Sequencing**

Illumina-based whole genome sequencing and assembly of strain PBC has been previously described (Gan et al., 2012b). To generate sequencing data using the MinION device, the gDNA of strain PBC was extracted from a 5-day old culture on nutrient agar using a slightly modified SDS-lysis method (Sokolov, 2000). Five micrograms of gDNA was subsequently used for library construction using the Nanopore sequencing kit SQK-NSK007 (Oxford Nanopore Technologies, Oxford, United Kingdom) according to the manufacturer’s instructions. The library was subsequently loaded onto an R9 MINION flowcell (Oxford Nanopore Technologies, Oxford, United Kingdom) and run for 48 h. 2D-basecalling was performed on the cloud-based Metrichor. Fast5 to fasta conversion used Nanocall (David et al., 2016).

**Hybrid Genome Assembly and Genome Annotation**

SPAdes version 3.8.1 (Antipov et al., 2016) was used for an initial hybrid genome assembly incorporating both Illumina
and Nanopore reads. Contigs longer than 1,000 bp were then selected for in silico scaffolding and gap-closing using npScarf (Cao et al., 2016). The assembled genome was annotated using the NCBI Prokaryotic annotation pipeline (Tatusova et al., 2016) and manually curated to include annotations for the pcaH2, pcaG2, pcaB2 and 4SLH genes coding for protocatechuate 3,4-dioxygenase alpha subunit type II, protocatechuate 3,4-dioxygenase beta subunit type II, 3-carboxymucononate cycloisomerase type II and 4-sulfomuconolactone hydrolase, respectively, that were missed by the default annotation setting. The identification and annotation of prophage sequences were performed using the PHASTER web-server (Arndt et al., 2016). Visualization of the complete genome of strain PBC and genomic comparisons with public available Hydrogenophaga genomes (Table 1) were performed using Blast Ring Image Generator (blastN e-value setting of 1e-10) (Alikhan et al., 2011). Mapping of Nanopore reads and Illumina contigs was performed using Minimap2 and subsequently visualized in Integrative Genomics Viewer version 3 (Thorvaldsdóttir et al., 2013). Generation of Illumina contigs was done by assembling Illumina-only reads using SPAdes version 3.8.1 and filtering for contigs with coverage and length of more than 5x and 300 bp, respectively (Antipov et al., 2016).

**In Silico Genome–Genome Hybridization**

Pair-wise average nucleotide identity of strain PBC against the publicly available genome sequences of Hydrogenophaga species were performed using JSpecies V1.2 (ANIm setting) (Richter et al., 2016). Strains exhibiting pair-wise ANI of more than 95% were considered as members of the same genospecies. A heatmap was plotted in R version 3 using the pheatmap library and 95% were considered as members of the same genospecies. Strains exhibiting pair-wise ANI of more than 5% were considered as members of the same genospecies.

**Whole Genome Phylogeny**

Whole proteome was predicted using Prodigal (-p meta setting) and piped into PhyloPhlan for the identification of conserved proteins (Hyatt et al., 2010; Segata et al., 2013). The concatenated alignment generated by PhyloPhlan was subsequently trimmed using trimAl version 1.9 (-automated1 setting) and used to construct a maximum likelihood tree using IQTree version 5.15 with 1000 ultrafast bootstraps (Capella-Gutíerrez et al., 2009; Nguyen et al., 2015). Visualization and annotation of trees was performed with MEGA6 (Tamura et al., 2013).

**Comparative Genomics Analysis**

HMMsearch version 3.1b was used for the identification of proteins of interest utilizing either the PFAM (−E 1e−5 setting) or TIGRFAM (−cut_nc or −E 1e−5 setting) HMM profiles (Haft et al., 2003; Eddy, 2011). Protein alignment, trimming, phylogenetic tree construction and visualization were performed using MAFFT-linsi, TrimAl, IQTree and FigTree, respectively (Capella-Gutiérrez et al., 2009; Katoh and Standley, 2013; Nguyen et al., 2015). Visualization of gene neighborhood was conducted using EasyFig (blastN setting) (Sullivan et al., 2011).

**RESULTS AND DISCUSSION**

**MinION Output and Hybrid Genome Assembly**

A total of 90,491 1D and 51,630 2D reads were generated from a full 48-h MinION run. After filtering for reads longer than 2,500 bp, 14,404 1D and 12,959 2D reads were retained for genome assembly. The final data output used for assembly was 173 megabases representing approximately 33x genome coverage of strain PBC. The average read length was 6,350 bp.

### Table 1: Hydrogenophaga strains used for comparative genomics analysis.

| Organism                          | Accession number | Isolation source                                      | Reference          |
|-----------------------------------|------------------|-------------------------------------------------------|--------------------|
| Hydrogenophaga intermedia PBC    | CP017311         | Textile waste water                                   | This study         |
| Hydrogenophaga intermedia S1 (DSM 5680<sup>T</sup>) | CCAE01         | Waste water with 4-aminobenzenesulphonate as sole source of carbon | Kim and Gan, 2017  |
| Hydrogenophaga sp. Root209        | LMIED1           | Root of Arabidopsis thaliana                          | Bai et al., 2015   |
| Hydrogenophaga pseudoflava NBRC 102511<sup>T</sup> | BCWO01         | River water                                           | unpublished        |
| Hydrogenophaga palleriini NBRC 102513<sup>T</sup> | BCTJ01         | Water enriched for hydrogen bacteria in an atmosphere containing 6% oxygen | unpublished        |
| Hydrogenophaga taeniospiralis NBRC 102512<sup>T</sup> | BCWR01         | Soil                                                  | unpublished        |
| Hydrogenophaga flavia NBRC 102514<sup>T</sup> | BCTF01         | Mud from ditch                                       | unpublished        |
| Hydrogenophaga sp. RAC07          | CP016449         | Algal phycosphere                                     | Fixen et al., 2016 |
| Hydrogenophaga sp. LPB0072        | LWDF01           | Pacific oyster (Crassostrea gigas)                    | Unpublished        |
| Hydrogenophaga sp. PML113         | MIYM01           | Contaminated water                                    | Unpublished        |
| Hydrogenophaga sp. IBVS2          | NFTU01           | Unknown                                               | Unpublished        |
| Hydrogenophaga sp. IBVS1          | NFUU01           | Unknown                                               | Unpublished        |
| Hydrogenophaga sp. A37            | MUNZ01           | Fjalar soil                                           | Unpublished        |
| Hydrogenophaga sp. H7             | MOIC01           | Coal mine                                             | Unpublished        |

<sup>1</sup>https://github.com/lh3/minimap2
and the longest read generated was 126,959 bp. Initial Spades hybrid assembly using both Illumina and MinION filtered reads generated 5 scaffolds (>10x coverage, >1,000 bp) which were subsequently scaffolded and gap-closed into a single contig using Npscarf. The complete genome of strain PBC is 5,232,477 bp in length with a GC content of 68.4%. Based on NCBI annotation pipeline, the complete genome consists of 4,904 protein-coding genes, 43 tRNA and single-copy 5S, 16S, and 23S rRNA genes. Comparisons of strain PBC and other Hydrogenophaga species revealed two genomic regions...
Hydrogenophaga trait of deaminate 4-aminobenzenesulfoate may be a defining metabolic trait of Hydrogenophaga genomes, suggesting that the ability to deaminate 4-aminobenzenesulfonate in strain PBC are clustered in two distinct genomic regions. Interestingly, the genes responsible for the catabolism of 4-aminobenzenesulfonate biodegradation in Hydrogenophaga strains (Figure 1A) have the highest number of homologous proteins to Enterobacter phage Arya (NCBI Reference Sequence: NC_0310480) isolated from a termite gut.

**New Genomic Insight into 4-Aminobenzenesulfonate Biodegradation in Hydrogenophaga**

Genes responsible for the catabolism of 4-aminobenzenesulfonate in strain PBC are clustered in two distinct genomic regions. Interestingly, the sad operon appears to be fairly conserved in all currently reported Hydrogenophaga genomes, suggesting that the ability to deaminate 4-aminobenzenesulfonate may be a defining metabolic trait of Hydrogenophaga strains. However, given the diverse isolation sources of Hydrogenophaga strains (Table 1) and their presumably limited exposure to xenobiotic compounds, it is likely that the original substrate of the 4-aminobenzenesulfonate dioxygenase is a naturally occurring compound with close structural homology to 4-aminobenzenesulfonate.

In contradistinction, the genomic region containing pcaH2G2, pcaB2 and 4-SLH involved in the conversion of 4-sulfocatechol to maleylacetate is conserved only in the Hydrogenophaga intermedia strains (Figure 1B). In addition, it is also worth noting that this region was identified as a phage region and flanked by genes coding for IS6 transposable elements (Figure 1C), an indication of the mobility of the pcaH2G2, pcaB2 and 4-SLH gene cluster presumably through phage transduction (Hickman et al., 2010). As expected, the presence of multiple transposase genes in this genomic region led to a fragmented initial Illumina read assembly (Figure 1C) due to the inability of short reads to resolve the repetitive regions (Phillippy, 2017). This shortcoming was overcome by integrating Nanopore long reads into the assembly as evidenced by their ability to span repeats in the genome (Figure 1C).

It is tempting to speculate that the acquisition of this gene cluster may be a key factor leading to complete 4-aminobenzenesulfonate biodegradation in Hydrogenophaga. Recently, the complete genome of Novosphingobium resinovorum SA1, another well-studied 4-aminobenzenesulfonate degrader, has been reported and comparison of this genomic region and that of the homologous region in Novosphingobium resinovorum SA1, revealed a comparable gene organization, e.g., pcaH2-pcaG2-tnp-pcaB2-4SLH-MAR in H. intermedia PBC (Figure 1C) and pcaB2 (scaA)-4SLH (scaB)-MAR(scaC)-oxidoreductase-pcaH2(scaE)-pcaG2(scaF) in Novosphingobium resinovorum SA1 (Hegedüs et al., 2017), suggesting recently shared ancestry of the currently known 4-sulfocatechol biodegradation pathway and possibly inter-class (alphaproteobacteria-betaproteobacteria) horizontal gene transfer of this gene cluster.

**H. pseudoflava and H. flava Belong to Two Genospecies Despite Their High Similarity in Genotypic and Protein Electrophoretic Profiles**

Hydrogenophaga flava was initially designated as the type species of the genus Hydrogenophaga but due to its slow and unreliable growth, H. pseudoflava has been proposed for use as an alternative reference taxon for the genus as it exhibits similar genotypic and protein electrophoretic profiles to that of H. flava (Willems et al., 1989). In silico genome-genome hybridization indicates that although H. flava and H. pseudoflava are closely related to one another, they are clearly two distinct genospecies with a pairwise ANIm of 93% (Figure 2). A more in-depth analysis of the H. flava genome may be useful to identify its additional nutritional requirements for optimal and consistent growth on culture medium. Such a genomic approach has been successfully demonstrated in Clostridium tyrobutyricum and other Clostridia associated with butyric acid fermentation leading to the identification of key vitamins and several amino acids essential for growth (Storari et al., 2016). Furthermore, in silico genome-genome hybridization analysis also indicates that H. sp. H7, which was previously isolated from a coal mine in China (Table 1), could be reclassified to the species H. pseudoflava given its high pairwise ANIm (97%) to H. pseudoflava NBRC 102511T (Figure 2).

**Genome-Based Phylogeny of the Family Comamonadaceae Supports Hydrogenophaga as a Monophyletic Group but Indicates Taxonomic Incongruence in the Genera Acidovorax and Comamonas**

Maximum likelihood tree based on 400 universal conserved proteins provides maximal support for the monophyletic clustering of all currently sequenced Hydrogenophaga strains with...
FIGURE 3 | Maximum likelihood phylogenomic tree of the family Comamonadaceae. The tree was rooted with members of the genus Burkholderia belonging to the family Burkholderiaceae, as the outgroup. Numbers in brackets indicate the number of branches collapsed. Asterisks indicate the presence of one member with inconsistent taxonomic labeling, e.g., different genus name within the collapsed branch. Numbers above branches indicate ultra-fast bootstrap support values. Table next to the Hydrogenophaga clade show the number of proteins showing significant hits to HMM profiles associated with bacterial hydrogenase system.
**H. intermedia** strains and *H. sp* PML 113 being basal to the rest of *Hydrogenophaga* (**Figure 3**). The genus *Hydrogenophaga* clade shares a sister relationship with a group of highly alkaliphilic hydrogen-utilizing Comamonadaceae bacteria isolated from an alkaline serpentinizing springs at The Cedars, California (Suzuki et al., 2014). Beyond the genus *Hydrogenophaga*, however, considerable taxonomic incongruence was observed in the genera *Acidovorax* and *Comamonas*. Some members of the *Acidovorax*, such as *Acidovorax* sp. strains 121606, 202149 and 12322-1, could be reclassified as members of the genus *Comamonas* based on their phylogenetic affinities. Within the genus *Comamonas*, *Comamonas badia* DSM 17552T (Suzuki et al., 2014) is peculiar as it does not cluster with a majority of other *Comamonas*, but instead forms a sister group with members of the genus *Alicycliphilus*, albeit with moderate bootstrap support. In addition, *Comamonas granuli* NBRC 101663T appears to be considerably divergent from other *Comamonas* species given its basal position in the clade containing all currently sequenced *Comamonas* strains (Suzuki et al., 2014). A more in-depth taxonomic investigation based on the percentage of conserved proteins (POCP) to specifically re-define the genus boundary between *Comamonas* and *Acidovorax* should be undertaken (Qin et al., 2014).

### Genomic Potential for Hydrogen Oxidation Is Restricted to a Few Members of Hydrogenophaga

The genus *Hydrogenophaga* was originally proposed by Willems et al. (1989) to group “yellow-pigmented hydrogen-oxidizing [Pseudomonas] species belonging to the acidovorans rRNA complex.” However, by identifying currently known HMM profiles related to hydrogenase system, genomic potential for hydrogen oxidation was only identified in 7 *Hydrogenophaga* strains of which 6 strains are closely related as shown by their monophyletic clustering in the phylogenomic tree (**Figure 3**). The 7 *Hydrogenophaga* strains include the original strains that were initially used to describe the genus *Hydrogenophaga*, such as *H. palleronii*, *H. taeniospralis*, *H. flava* and *H. pseudoflava*, corroborating their previously demonstrated in vitro hydrogen oxidizing ability (Willems et al., 1989; Contzen et al., 2000). The reported inability of *H. intermedia* S1T and *H. atypica* DSM 15342T to oxidize hydrogen in vitro (Contzen et al., 2000; Kampfer et al., 2005) correlates with the absence of key genes associated with hydrogen metabolism (**Figure 3**). However, the absence of genomic potential for hydrogen catabolism in nearly half of the currently sequenced *Hydrogenophaga* strains is unexpected and indicates that the description genus of *Hydrogenophaga* as “hydrogen eating bacteria” is not sustainable and warrants revision.

### p-Aminobenzoic Acid Auxotrophy within Hydrogenophaga

Mapping of *Hydrogenophaga* genome sequences to the contig of *H. palleronii* NBRC102513 (BC701000048, 24 kbp) containing *pabB* revealed that only 8 out of 14 genomes exhibit nucleotide similarity (>40% identity and E-value < 1E-5) to the *pabB* gene region (**Figure 4A**). Of the eight genomes, only partial *pabB* hits were observed for strains H7, NBRC102514, Root209, NBRC102511 and A37, hinting divergence at the nucleotide level. Interestingly, only two proteins belonging to *H. palleronii* and *H. sp*. PML113 achieved an HMMscore that is above the domain noise cutoff score for bona fide PabB (TIGR00553) (**Supplementary Figure S1**). On the contrary, strain IBVS2 despite exhibiting nucleotide similarity to the *pabB* gene region, does not code for protein with HMMscore above the TIGR00553 noise cutoff. However, it is worth noting that multiple proteins with HMMscore slightly below the TIGR00553 noise cutoff could be identified from the annotated *Hydrogenophaga* genomes (**Supplementary Figure S1**). These proteins appear to be more closely related to PabB than TrpE as evidenced by their extremely low HMMscore to TrpE (TIGR00564) in comparison with other putative TrpE proteins and their phylogenetic affiliation to the PabB clade (**Supplementary Figure S2 and Data Sheet 1**). TrpE is involved in the synthesis of anthranilate from chorismate and ammonia. Functional characterization of these divergent PabB homologs will be useful to explore their role in pABA synthesis as of them belong to *Hydrogenophaga pseudoflava* that has been shown to grow on minimal medium without pABA supplementation (Povolo et al., 2013).

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**FIGURE 4** | Prevalence of *pabB* among *Hydrogenophaga* strains based on genomic and phylogenetic analyses. (A) Homology analysis against the *pabB*-containing genomic region of *Hydrogenophaga palleronii* **(B)** Easyfip visualization of homologous genomic regions containing *pabB* between *H. intermedia* PBC and *H. palleronii*. Shaded regions indicate regions with E-value score of less than 1e-5 with the percentage of nucleotide identity expressed along a gray scale. Hypothetical proteins are indicated by arrows containing black circles.
The absence pabB in *H. intermedia* strains PBC and S1^T^ is consistent with their previously reported reliance on either external nutrient supplementation or helper strain for pABA (Dangmann et al., 1996; Gan et al., 2011b; Kim and Gan, 2017). A recent genomic analysis of *Hydrogenophaga intermedia* PBC helper strain, *Ralstonia* sp. PBA, showed the presence of two PabB proteins of which one of them was fused to PabC (Gan et al., 2012a; Kim and Gan, 2017), potentially enabling the overproduction of pABA to sustain the growth of both strains when they were co-cultured in vitamin-free minimal medium containing 4-aminobenzenesulfonate as the sole carbon and nitrogen source. A closer comparison of the genomic region containing genes spatially associated with pabB in *H. intermedia* PBC and *H. palleronii* NBRC102513 suggests that pabB may be lost through gene deletion (Figure 4B). Surprisingly, strains LPB0072, RAC07, Root209 and BVHSV1 appear to also lack the genomic potential for pABA synthesis. A future study investigating the ability of these strains to grow on defined medium without pABA supplementation will be necessary to verify their predicted pABA auxotrophy as most *Hydrogenophaga* strains were isolated on nutrient rich medium, which will contain traced amount of pABA. For example, strains Root209 and RAC07 were maintained on tryptic soy agar and minimal medium with yeast extract supplementation, respectively (Bai et al., 2015; Fixen et al., 2016).

**DATA ACCESS**

The complete genome of *Hydrogenophaga intermedia* PBC has been deposited under the accession number CP017311. Raw data for both Illumina and Nanopore data have been placed under the SRA under the SRA project ID SRP092076.

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**AUTHOR CONTRIBUTIONS**

HG performed gDNA extraction, and bioinformatics analysis and drafted the manuscript. YL performed library preparation and MINION sequencing. CA supervised the study.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fmicb.2017.01880/full#supplementary-material

**FIGURE S1** | Sequence scores of *Hydrogenophaga* proteins with significant hits (E-value < 10^-5) to TIGR00564 or TIGR00553. Dotted lines represent the TIGRFAM model noise cutoff bit score thresholds.

**FIGURE S2** | Maximum likelihood tree of TrpE and PabB. Node values indicate ultrafast bootstrap support values. Red branches indicate proteins with high sequence scores to TIGR00564 corresponding to bona fide TrpE. Blue and green branches indicate proteins with low scores to TIGR00564 but high scores to TIGR00553 (bona fide PabB). In addition, proteins associated with blue branches exhibit scores that are either higher than or very close to the noise cutoff score TIGR00553.

**DATA SHEET 1** | *Hydrogenophaga* protein sequences used for the construction of maximum likelihood tree.
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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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