

# Design of a degenerate primer pair to target a bacterial functional community: The hppd bacterial gene coding for the enzyme targeted by herbicides, a study case

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1	Design of a degenerate primer pair to target a bacterial functional community:		
2	the <i>hppd</i> bacterial gene coding for the enzyme targeted by herbicides, a study case		
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10	Abstract: The present work aimed to design a degenerate primer pair to target a large part o		
11	the hppd soil bacterial community, possibly affected by herbicides. We validated these		
12	primers by qPCR and high-throughput sequencing analysis of soil samples.		
13	Keywords: hppd, degenerate primers, soil bacterial community.		
14	Highlights:		
15	• A novel way to design a degenerate primer pair targeting a functional community		
16	• Soil bacterial <i>hppd</i> community abundance and diversity can be monitored		
17	• The effects of HRAC F2 herbicides on soil communities can now be assessed		
18	In recent scientific opinions, the European Food Safety Authority proposed to include soil		
19	ecosystemic services as a specific protection goal for risk assessment of pesticides and to		
20	protect soil microorganisms at the level of functional groups (EFSA, 2010). Another scientific		
21	opinion underlines the lack of standardized methods to measure the effects of pesticides on		
22	structural and functional endpoints of the microbial community (EFSA, 2017). To fill this		
23	knowledge gap, microbial genes encoding enzymes inhibited by substances with herbicidal		
24	activity could be ideal biomarkers of the direct effects of these molecules on soi		
25	microorganisms (Thiour-Mauprivez et al., 2019).		

Screening of these active substances according to their mode of action led to the 26 identification of those that affect the cell metabolism. A list of targeted enzymes was drawn 27 up by the HRAC (https://hracglobal.com/tools/world-of-herbicides-map). Among them, 4-28 hydroxyphenylpyruvate dioxygenase (4-HPPD; EC 1.13.11.27) appeared as a possible 29 biomarker sensitive to the herbicides of the HRAC (Herbicide Resistance Action Committee) 30 F2 group (list of 13 active substances with herbicidal activity, supplementary file 1). This 31 enzyme transforms tyrosine into fumarate and acetoacetate. These two compounds enter the 32 central metabolism and lead to the formation of pyomelanine, a brown pigment that has 33 antioxidant properties and protects fungi and bacteria from reactive oxygen species (ROS) 34 (Langfelder et al., 2003; Turick et al., 2009). To our best knowledge, 2,869 HPPD protein 35 sequences from various bacterial species are available in the Pfam database 36 (http://pfam.xfam.org/family/PF14696) (figure 1). One could hypothesize that among the 37 38 numerous populations harbouring 4-HPPD within the bacterial community, several of them might be sensitive to HRAC F2 herbicides. From this point of view, both a priori and a 39 40 posteriori risk assessment of HRAC F2 herbicides on soil microorganisms might be assessed by monitoring parameters of the *hppd* bacterial community. 41

In this context, the aim of this study was to develop a molecular tool to monitor the 42 abundance and diversity of this community in the soil. In order to target as many bacterial 43 hppd sequences as possible, we developed a pipeline (supplementary file 2) to design a 44 degenerate primer pair targeting a broad spectrum of bacterial 4-HPPD genes. This was 45 challenging since the known bacterial 4-HPPD proteins available on Pfam presented both 46 length and sequence polymorphisms. Sequence lengths varied from 237 to 491 amino acids 47 (95% of the 4-HPPD sequences were 325 to 360 amino acids in length), and the most distant 48 49 protein sequences showed only 22% similarity (79/354 identical amino acids). The corresponding nucleic acid sequences were retrieved from GenBank, aligned with Muscle 50

(Edgar, 2004), and too long, too short or too different sequences were removed manually. The 51 2,506 remaining sequences were converted into protein sequences using Mega7 (Kumar et al., 52 2016), resulting in a clean bacterial 4-HPPD database (see taxonomic affiliation of these 53 sequences in table 1). The latter was submitted to j-CODEHOP (Boyce et al., 2009) to 54 identify conserved motifs candidate for primer design. Two motifs were identified: HIA 55 (harboured by 1,957 out of 2,506 sequences) and FFE (found in 2,361 sequences) (in pink and 56 violet in **figure 1**). Twenty-base degenerate primers were then designed so that the nine bases 57 in 3'-terminal position perfectly annealed the HIA or FFE motifs shared by 1,812 of the 2,506 58 analysed sequences (figure 2). 59

To validate our primer pair, three soil samples were collected from the surface layer (0-20 60 cm) of an arable field (University of Perpignan, France) (see properties in supplementary file 61 3). Soil samples were prepared as previously described (Romdhane et al., 2016). Nucleic 62 63 acids were extracted with an RNeasy PowerSoil DNA Elution Kit (Qiagen) and the hppd gene was successfully amplified by PCR from these DNA extracts as described in supplementary 64 65 file 4. The affiliation of the amplicons to the *hppd* gene sequence was checked by cloning and sequencing. Ninety-two percent of the 24 sequenced amplicons were related to hppd 66 sequences (sequences submitted and released upon manuscript acceptance). The hppd 67 community relative abundance was assessed by qPCR as described before (Romdhane et al., 68 2016) except for the HIA-F/FFE-R primer pair which was used at 10µM in our final reaction 69 mixture with an annealing temperature of 54°C. A calibration curve was generated in 70 triplicate using serial dilutions of the standard ranging from  $10^2$  to  $10^7$  copies *per* reaction. 71 72 Three no-template controls were included for each qPCR assay. The calibration curve showed an acceptable efficiency of 79% ( $R^2 = 0.99$ ). The detection limit was around 10<sup>2</sup> copies of 73 74 hppd copies per reaction (data not shown). Relative abundance was 421±43 hppd sequences

*per* 1,000 sequences of 16S rDNA. *In silico* observations confirmed that *hppd* gene sequences
were quite abundant among the soil bacterial community.

hppd sequence diversity of the amplicons obtained with a two-step PCR was assessed by 77 Illumina 2\*250 base pair MiSeq sequencing run (BioEnvironnement, University of Perpignan, 78 France). The first PCR involved 27 cycles of amplification. It was performed in duplicate for 79 each of the three soil DNA samples, and one no-template control using the NGS HIA F (5'-80  ${\tt TCGTCGGCAGCGTCAGATGTGTATAAGAGACAGNNNNARGGBATYCAGCAYATY}$ 81 GCN-3') NGS\_FFE\_R (5'-82 and GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAGNNNNCGYTGDATRAWYTCRA 83 ARAA-3') primer pair with a specific tail added to match the second PCR primer pair. Six 84 microliters of the pooled first PCR products were used as a template for the second PCR that 85 involved eight cycles of amplification with barcoded primers in duplicate. After pooling and 86 87 amplicon size verification by electrophoresis, hppd amplicons were normalized using a SequalPrep<sup>TM</sup> Normalization Plate Kit (Invitrogen). Ten microliters of each sample were used 88 89 for Illumina MiSeq sequencing. After assembly using PEAR (Zhang et al., 2014) and quality 90 check using the QIIME pipeline (Caporaso et al., 2010), a total of 744,084 hppd nucleic sequences were obtained.. These sequences were corrected and converted into proteins using 91 framebot (Wang et al., 2013), then clustered using cd-hit (Li and Godzik, 2006) (threshold: 92 100%). The resulting 62,409 clusters were compared to the 4-HPPD database using blast, and 93 affiliated to the closest relative sequence (minimum similarity: 57%). The designed primers 94 allowed us to retrieve hppd sequences affiliated to most phylogenetic groups from our 95 96 database, as shown in figure 1. However, only a few sequences related to Actinobacteria were obtained. This might be due not only to primer efficiency regarding this group but also to 97 98 various biases such as DNA extraction and/or low abundance of Gram-positive bacteria in this

soil. A small number of *hppd* sequences that contained only one (5.6%) or no motif (2.7%)

100 were also amplified, suggesting that they may correspond to sequences not yet described.

101 This primer pair appears to be suitable for measuring the abundance of the *hppd* bacterial 102 community and estimating its diversity in an arable soil. Further studies will aim to monitor 103 the evolution of the *hppd* bacterial community exposed to herbicides of the HRAC F2 group. 104 It is noteworthy that this primer design method could also be applied to amplify other

105 functional genes, paving the way for the analysis of other bacterial subcommunities.

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### 176 **Figure legends**

**Figure 1.** Phylogenetic tree of the 4-HPPD database derived from Pfam, constructed with FastTree (Price *et al.*, 2009) (Neighbour-Joining) and visualized with iTol (Letunic and Bork, 2019). The inner circle represents the affiliation of the 4-HPPD sequences to the main microbial groups indicated by different colours. The middle circle depicts the presence of the HIA and/or FFE motifs in the HPPD sequences, indicated by different colours. The outer circle shows the affiliation of the OTUs found in Perpignan soil samples to known HPPD sequences.

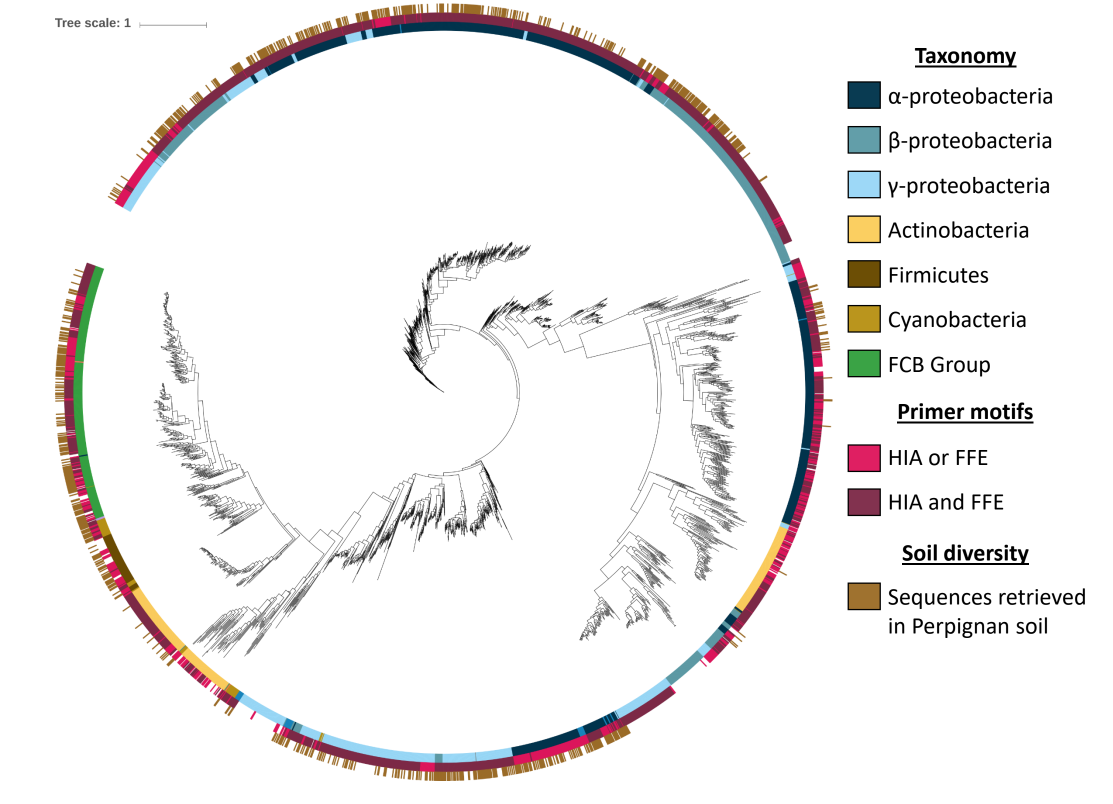
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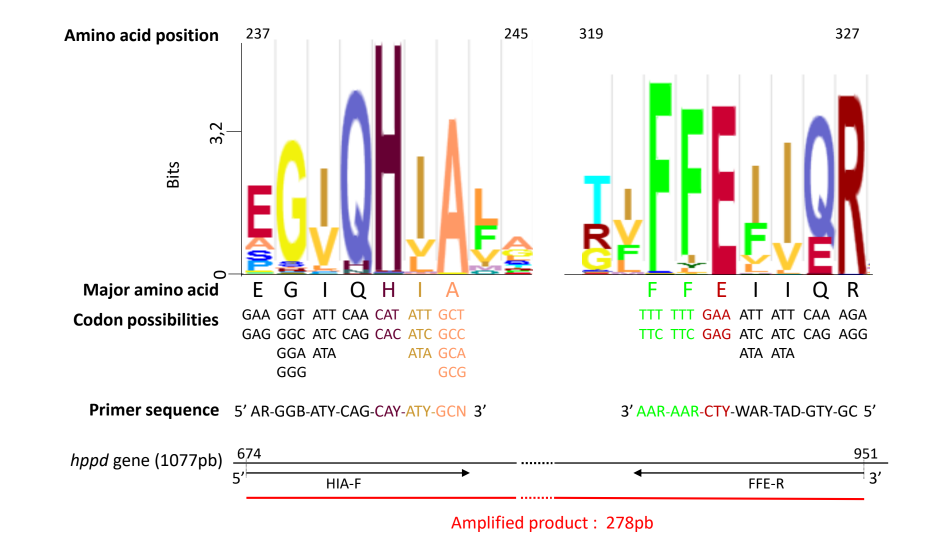
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Figure 2. HMM logos constructed with Skylign (Wheeler *et al.*, 2014) showing the
occurrence of the amino acids which composed our primer pair. R, A or G; B, C or G or T; Y,
C or T; N, any base; D, A or G or T; W, A or T. The amino acid position and the amplified
product length were determined in the *hppd* sequence of *Pseudomonas fluroescens* F113
(NC\_016830.1).

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**Figure 1.** Phylogenetic tree of the 4-HPPD database derived from Pfam, constructed with FastTree<sup>12</sup> (Neighbor-Joining) and visualized with iTol<sup>13</sup>.



**Figure 2.** HMM logos constructed with Skylign<sup>14</sup> showing the occurrence of the amino acids which composed our primer pair.

Phylum	Class	Number of sequences
Proteobacteria	α-proteobacteria	822
	γ-proteobacteria	529
	β-proteobacteria	484
Terrabacteria group	Actinobacteria	265
	Firmicutes	67
	Cyanobacteria	37
FCB group	Bacteroidetes	298
	Gemmatimonadetes	3
	Acidobacteria	1