The genes and enzymes of phosphonate metabolism by bacteria, and their distribution in the marine environment

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Phosphonates are compounds that contain the chemically stable carbon–phosphorus (C–P) bond. They are widely distributed amongst more primitive life forms including many marine invertebrates and constitute a significant component of the dissolved organic phosphorus reservoir in the oceans. Virtually all biogenic C–P compounds are synthesized by a pathway in which the key step is the intramolecular rearrangement of phosphoenolpyruvate to phosphopyruvate. However C–P bond cleavage by degradative microorganisms is catalyzed by a number of enzymes – C–P lyases, C–P hydrolases, and others of as-yet-uncharacterized mechanism. Expression of some of the pathways of phosphonate catabolism is controlled by ambient levels of inorganic P (Pi) but for others it is Pi-independent. In this report we review the enzymology of C–P bond metabolism in bacteria, and also present the results of an in silico investigation of the distribution of the genes that encode the pathways responsible, in both bacterial genomes and in marine metagenomic libraries, and their likely modes of regulation. Interrogation of currently available whole-genome bacterial sequences indicates that some 10% contain genes encoding putative pathways of phosphonate biosynthesis while ∼40% encode one or more pathways of phosphonate catabolism. Analysis of metagenomic data from the global ocean survey suggests that some 10 and 30%, respectively, of bacterial genomes across the sites sampled encode these pathways. Catabolic routes involving phosphonoacetate hydrolase, C–P lyase(s), and an uncharacterized 2-aminoethylphosphonate degradative sequence were predominant, and it is likely that both substrate-inducible and Pi-repressible mechanisms are involved in their regulation. The data we present indicate the likely importance of phosphonate-P in global biogeochemical P cycling, and by extension its role in marine productivity and in carbon and nitrogen dynamics in the oceans.

Keywords: phosphonate, C–P bond, marine bacteria, phosphorus cycling

PHOSPHORUS IN THE OCEANS

Phosphorus (P) is usually found in the biosphere in its highest oxidation state (+5) as orthophosphate (Pi), which is thought to be its most bioavailable form. P availability has long been identified as a key determinant of marine productivity (Dyhrman and Haley, 2006; Paytan and McLaughlin, 2007; Dyhrman et al., 2009) and its cycling is closely linked to carbon (C) and nitrogen (N) dynamics in the oceans. In oligotrophic regions, and seasonally in the surface waters of a broad range of marine environments, P concentrations can be extremely low, e.g., most of the central oceanic gyres have levels close to the limit of chemical detection (<1 nM). In these instances recent climatically influenced extension of the N-fixation season appears to have led to a shift from an N- to a P-limited ecosystem (Karl et al., 2001). Correlated with this has been an increase in levels of Dissolved Organic Phosphorus (DOP), relative to those of P., Several recent studies suggest that DOP is a significant additional P source that is available to the marine microbiota in a variety of trophic regimes (Björkman and Karl, 2003; Luo et al., 2009, 2011; Martinez et al., 2010). For example a study in the oligotrophic North Pacific subtropical gyre showed that DOP accounted for up to 80% of total soluble P and was rapidly utilized by the microbial fraction (Björkman and Karl, 2003). It may thus play an important role in determining the composition and community dynamics of marine microbial populations.

PHOSPHONATES IN THE MARINE ENVIRONMENT

An important component of marine DOP is phosphonate-P, which occurs widely in nature in a range of biogenic molecules that contain the highly stable and apparently ancient direct carbon–phosphorus (C–P) bond, rather than the more common C–O–P bond found in phosphate esters (Ternan et al., 1998; Quinn et al., 2007; White and Metcalf, 2007). Phosphonates are known to be produced by many primitive life forms, including a wide range of marine invertebrates in which phosphonate-P may comprise as much as 50% of the total P content (Quin, 1965, 2000). The presence of phosphonates at levels of some 10% of total P in cells of the bloom-forming marine cyanobacterium Tri chodesmium erythraeum IMS 101 has also recently been demonstrated (Dyhrman et al., 2009). Natural-product phosphonates such as 2-aminoethylphosphonic acid (2-AEP; ciliatine) or 2-amino-3-phosphonopropionic acid (phosphoanalanine) are normally found as side groups on exopolysaccharides or glycoproteins,
or in the polar head groups of membrane phospholipids (Hilderbrand, 1983); their presence may increase structural rigidity or protect against enzymatic degradation, since the C–P bond is resistant to the action of phosphatases. However a growing range of small-molecule phosphonates are also known to be produced (reviewed by Metcalf and van der Donk, 2009); many are likely to serve as antimetabolites (e.g., fosfomycin, fosmidomycin, alafosfalin, and phosphonothricin) due to their structural analogies to phosphate esters or carboxylic acids.

The qualitative composition and quantitative importance of phosphate-P in the oceans have been difficult to establish definitively because of analytical limitations. Nonetheless studies using 31P-NMR indicate that C–P compounds comprise more than 25% of ultrafiltered DOM, and 23% of total particulate P (Clark et al., 1998; Kolowith et al., 2001). *In situ* evidence for high rates of phosphonate remineralization has also been presented, for example by Benitez-Nelson et al. (2004). Moreover a number of recent field, laboratory, and genome-enabled studies (Dyhrman et al., 2006; Quinn et al., 2006; Karl et al., 2008; Gilbert et al., 2009; Ilikchyan et al., 2009; Martinez et al., 2010; Thomas et al., 2010; Luo et al., 2011) also suggest that phosphonate-P may constitute a significant P source in the oceans. Additionally – if circumstantial – evidence for this view is provided by the fact that the “streamlined” genome of SAR 11, the most abundant bacterial clade on earth, contains (and expresses *in situ*) phosphonate transport genes (Sowell et al., 2009). Despite this growing body of information, however, our understanding of the factors that influence the production and consumption of phosphonates in the oceans remains limited (Dyhrman et al., 2007, 2009).

**BIOSYNTHESIS OF PHOSPHONATES**

Virtually all known pathways of phosphonate biosynthesis share a central C–P bond-forming reaction in which an intramolecular rearrangement of the intermediary metabolite phosphoenolpyruvate (PEP) is catalyzed by the enzyme PEP phosphomutase (Ppm; EC 5.4.2.9), with phosphoryl group transfer leading to formation of phosphonopyruvate (Bowman et al., 1988; Seidel et al., 1988; Metcalf and van der Donk, 2009; Figure 1A). Since the equilibrium of the reaction carried out by PEP phosphomutase strongly favors formation of the C-O-P bond of PEP, however, most C–P bond-synthesizing microorganisms contain a coupled enzyme, phosphonopyruvate decarboxylase (Ppd; EC 4.1.1.82), which subsequently converts phosphonopyruvate to phosphonoacetaldehyde in an irreversible reaction (Zhang et al., 2003; Figure 1A). Phosphonoacetaldehyde then forms the central metabolite in a range of subsequent biosynthetic sequences (Metcalf and van der Donk, 2009); for example in the biosynthesis of 2-AEP it is transaminated by the enzyme 2-AEP-pyruvate transaminase (EC 2.6.1.37; Barry et al., 1988).

**ENZYMES OF PHOSPHONATE CATABOLISM IN MICROORGANISMS**

The importance of phosphonates as a P reservoir in the global ocean is by now well-established but information on the molecular strategies employed by marine microorganisms for their utilization remains incomplete. Current understanding of the catabolic activities involved is summarized below.

**C–P LYASE**

The predominant route for microbial utilization of phosphonates has been thought to be the “C–P lyase” pathway (Ternan et al., 1998) and its presence and activity in marine microorganisms has recently been demonstrated (e.g., Dyhrman et al., 2006). However the broad-specificity C–P lyase multienzyme complex has not yet been fully characterized (Hove-Jensen et al., 2011) although a major technical obstacle to its elucidation has very recently been overcome with the first report (Kamat et al., 2011) of the *in vitro* reconstitution of the pathway in *Escherichia coli* and identification of the reaction intermediates. In *E. coli* the C–P lyase complex is comprised of 14 gene products which together catalyze the difficult dephosphonation reaction in a range of structurally diverse phosphonates which may contain either activated or unactivated C–P bonds (Wackett et al., 1987; Yakovleva et al., 1998). In each case the product is the corresponding hydrocarbon and Pi (Figure 1B).

Of these 14 genes, *phnCDE* are thought to encode an ATP-binding cassette transporter, and *phnF* a repressor protein, while the products of *phnNOP* are thought to have regulatory or other accessory roles (Hove-Jensen et al., 2010, 2011). Seven gene products (*PhnGHIJKLM*) constitute the “core” components of the C–P lyase reaction pathway, with *PhnJ* now known to catalyze its central reaction, the S-adenosyl-l-methionine-dependent radical cleavage of 5-phosphoribosyl-1-phosphate to produce 5-phosphoribosyl-1,2-cyclic phosphate and the corresponding alkane (Hove-Jensen et al., 2011; Jochimsen et al., 2011; Kamat et al., 2011). Variant forms of the C–P lyase enzyme complex with different substrate ranges are also known to exist (e.g., Kertesz et al., 1991; Sviridov et al., 2012), and other genes of as-yet unknown function have frequently been found associated with the C–P lyase operon (see Occurrence of the *phnJ* Component of the C–P Lyase Operon in Sequenced Bacterial Genomes). In addition the role of *PhnK* remains unknown (Kamat et al., 2011).

In all bacteria studied to date the genes that encode the C–P lyase pathway(s) have been found to be arranged in tandem and transcribed from a single promoter in an operon that is a member of the Pho regulon – a suite of genes whose products include proteins required for Pi-scavenging, and for the catabolism of alternative P sources. Expression of the genes of the Pho regulon is induced under conditions of Pi-starvation but is Pi-repressible (Quinn et al., 2007). Thus in *T. erythraeum* the genes of the C–P lyase pathway were expressed *in situ* at Pi concentrations of ~6 nmol l⁻¹, but not under Pi-replete conditions (Dyhrman et al., 2006). Similarly the aerobic production of methane from methylphosphonate by mixed marine microbial communities (likely to be mediated by C–P lyase) was demonstrated under Pi-stressed conditions (Karl et al., 2008).

All C–P bond cleavage enzymes had originally been thought to be similarly subject to Pho regulon control; however, it seems likely that in this event phosphonates could be utilized by microorganisms only as a source of P, since any Pi released during their catabolism in excess of that required for balanced cellular growth would repress any further degradation (McGrath et al., 1997; Quinn et al., 2007). Indeed, further phosphonate uptake by cells would also be repressed because the relevant high-affinity transporters are also Pho-regulated. However field and laboratory observations (Benitez-Nelson et al., 2004; Beversdorf et al., 2010) suggest that
**FIGURE 1 | Pathways for the microbial metabolism of phosphonates:**

(A) Phosphonate biosynthesis by phosphoenolpyruvate phosphomutase [1] and phosphonopyruvate decarboxylase [2].

(B) C–P lyase [3] pathway.

(C) 2-AEP catabolism by 2-AEP transaminase [4] and phosphonoacetaldehyde hydrolase [5].

(D) Phosphonoacetate catabolism by phosphonoacetate hydrolase [6].

(E) 2-AEP catabolism by 2-AEP transaminase [4], phosphonoacetaldehyde dehydrogenase [7], and phosphonoacetate hydrolase [6].

(F) Phosphoalanine catabolism by phosphoalanine transaminase [8] and phosphonopyruvate hydrolase [9].
phosphonate degradation in the oceans continues under conditions of Pi sufficiency. And indeed three inducible enzymes capable of C–P bond cleavage have been identified and characterized (Quinn et al., 2007). Unlike C–P lyase, which is capable of the cleavage of both activated and unactivated C–P bonds, each of these phosphohydrolases is uniquely specific to a natural-product phosphonate that contains an electron-withdrawing carbonyl group in the beta position to the P atom. Expression of all three enzymes has been shown to be Pi-insensitive and hence independent of Pho regulation — thus they are likely to play a role in the marine environment that is complementary to that of C–P lyase.

**PHOSPHONOACETALDEHYDE HYDROLASE**

The **phnX** gene product phosphonoacetaldehyde hydrolase (EC 3.11.1.1; phosphonatase) cleaves the C–P bond of its unique substrate to yield acetaldehyde and Pi (LaNauze et al., 1977; Dumora et al., 1989); it is encoded within a degradative operon for the mineralization of 2-aminoethylphosphonic acid (2-AEP, ciliatine). 2-AEP replaces its structural analog ethanolamine phosphate as a component of the phosphoglycerolipids common in the cell membrane of many invertebrates and is considered — along with its N-methylated derivatives — to be the most abundant and ubiquitous marine C–P compound (Hilderbrand, 1983; Hori et al., 1984; Quin and Quin, 2001). The mechanism of C–P bond cleavage by phosphonatase, a member of the halo acid dehalogenase superfamily, involves the formation of a Schiff base intermediate between a lysine residue at the active site of the enzyme and the phosphonoacetaldehyde carbonyl group; this activates the phosphonate group for attack by an active site nucleophile (Baker et al., 2011) which has also been designated as PhnY, both here and elsewhere in the literature. The PhnX enzyme is a member of the halo acid dehalogenase superfamily, involves the formation of a Schiff base intermediate between a lysine residue at the active site of the enzyme and the phosphonoacetaldehyde carbonyl group; this activates the phosphonate group for attack by an active site nucleophile (Baker et al., 2011; Chen et al., 2006). The PhnX enzyme is a member of the halo acid dehalogenase superfamily, involves the formation of a Schiff base intermediate between a lysine residue at the active site of the enzyme and the phosphonoacetaldehyde carbonyl group; this activates the phosphonate group for attack by an active site nucleophile (Baker et al., 2011; Chen et al., 2006).

**PHOSPHONOPYRUVATE HYDROLASE**

This enzyme was identified by Ternan and Quin (1998b) in cells of a *Burkholderia* strain that utilized phosphonoalanine (2-amino-3-phosphonopropionic acid) as sole C, N, and P source. Phosphonoalanine, the phosphonate analog of aspartic acid, is formed by the transamination of phosphonopyruvate (Horigane et al., 1989). It is a constituent of membrane phosphonolipids and is widely distributed amongst marine invertebrates (Kittredge and Hughes, 1964); its degradative pathway involves an initial transamination to phosphonopyruvate, followed by cleavage of the C–P bond by a metal-dependent phosphonopyruvate to yield pyruvate and Pi (Figure 1E). Phosphonopyruvate hydrolase (EC 3.11.1.3) is a member of the phosphonopyruvate phosphomutase/isoctirate lyase superfamily and was shown in *Variovorax* sp. Pa12 to be encoded within an operon that also contains phosphonoalanine transaminase and phosphonate transporter genes; its expression is regulated by an adjacent phosphonoalanine- and phosphonopyruvate-responsive LTRR (Kulakova et al., 2003, 2009). Phosphonopyruvate hydrolase shares 41% sequence identity with the C–P bond-forming enzyme, phosphonopyruvate phosphomutase (see Biosynthesis of Phosphonates above); a single amino acid substitution at the active site is likely to be responsible for the functional divergence of the two enzymes (Chen et al., 2006).

**OTHER ENZYMES OF PHOSPHONATE CATABOLISM**

There is also evidence for the existence of a number of less well-characterized C–P bond cleavage activities that are likely to be of significance in marine environments:

(a) Martinez et al. (2010) have demonstrated by functional gene complementation that a fragment of marine genomic DNA that encodes a putative 2-oxoglutarate dioxygenase (PhnY*) and a phosphohydrolase (PhnZ) conferred on *E. coli* the ability to utilize 2-AEP as a sole P source; however the biochemical reactions catalyzed by these genes remain uncharacterized. It will be important to avoid future confusion between the activity encoded by *phnY* and the unrelated phosphonoacetaldelyde dehydrogenase (see Phosphonoacetate Hydrolase above) recently described in *Sinorhizobium meliloti* by Borisova et al. (2011) which has also been designated as PhnY, both here and in a number of annotated genomes.

(b) McGrath et al. (1998) have reported the utilization of the phosphonate antibiotic fosfomycin (2-epoxypropylphosphonic acid) as the sole C and P source by a *Sinorhizobium huakuii* isolate. The catabolic reaction sequence involved has been further elucidated (McGrath et al.,
2009, 2011), and although the enzyme responsible for the C–P bond cleavage step has not yet been fully described its substrate is likely to be (R)-1-hydroxy-2-oxopropylphosphonic acid. Given that S. huakuii cells release essentially quantitative levels of Pi when using fosfomycin as a sole C source, expression of the C–P cleavage enzyme is clearly Pho-independent. (c) Mendz et al. (2005) and Ford et al. (2010) have presented evidence for the existence of novel, as-yet-undescribed pathways of Pi-insensitive phosphonate catabolism in Campylobacter and Helicobacter spp. The existence of further uncharacterized, but likely unrelated, C–P cleavage activities has also been reported in cells of Synechococcus spp. (Gomez-Garcia et al., 2011).

**DISTRIBUTION OF THE GENES OF PHOSPHONATE METABOLISM AMONGST SEQUENCED BACTERIA**

BLAST analysis of the occurrence of the genes of phosphonate metabolism amongst the bacterial genome sequences deposited in the NCBI GenBank database as at the beginning of 2011 was carried out. The query sequences from deduced peptides of confirmed function used to interrogate the database are listed in Table 1. A minimum expectation value of 1 × 10⁻¹⁰ and a minimum of 10% sequence identity over the entire sequence length were employed as criteria for homolog selection. In addition each gene identity was confirmed by inspection of its genomic context.

Out of the 1384 sequenced genomes in the database, 664 (48%) contained homologs that indicated possession of one or more of the known pathways of phosphonate metabolism (Table 2); these included 121 of the 190 marine bacterial genomes in the database. Fourfold more strains contained known genes associated with phosphonate catabolism than with C–P bond formation. Of the putative phosphonate degradative pathways detected, those involving C–P lyase and phosphonooacetaldehyde hydrolase comprised some 83% of the total. The abundance of *phnX* homologs encoding the substrate-specific phosphonooacetaldehyde hydrolase, and of homologs that encode the PhnD transporter which shows highest affinity for 2-AEP (Rizk et al., 2006) suggests that 2-AEP is likely to have been an important source of P through a long period of evolutionary history. An analysis of the distribution of each of the pathways and of the variation in their associated operons is presented below.

**OCCURRENCE IN SEQUENCED BACTERIAL GENOMES OF THE PHOSPHENOLOPYRUVATE PHOSPHOMUTASE/PHOSPHONOPYRUVATE DECARBOXYLASE OPERON FOR PHOSPHONATE BIOSYNTHESIS**

129 sequenced strains (almost 10% of the genomes contained in the NCBI GenBank database) contained the ppd gene indicative of the PEP phosphomutase/phosphonopyruvate decarboxylase pathway of phosphonate biosynthesis; seven of these were of marine origin. Of the 129 strains, 48 also possessed a pathway for phosphonate degradation, and 10 the associated *phnCD* operon encoding a phosphonate transporter linked to phosphonate catabolism (Huang et al., 2005). The phosphonate biosynthetic pathway was distributed amongst distant bacterial groups, but predominantly within the members of the proteobacteria, actinobacteria, and clostridia (Figure 2). Analysis of their *ppm/ppd* operon structures failed to reveal any clearly conserved gene composition, which may reflect the wide range of phosphonate metabolites that are produced by microorganisms (Circello et al., 2010; Johannes et al., 2010; Lee et al., 2010). Nevertheless, two enzymes frequently found to be encoded within the operon (in addition to PEP phosphomutase and phosphonopyruvate decarboxylase) were 2-AEP aminotransferase (PhnW) – observed in 60 strains – which plays a key role in the production of 2-AEP, and an acetalddehyde dehydrogenase (found in 16 strains), which shares strong homology (44% identity, 60% similarity) with the phosphonoacetalddehyde dehydrogenase (PhnY) known to be involved in the production of phosphonoacetate from phosphonoacetalddehyde and found in the phosphonoacetate hydrolase pathway of 2-AEP degradation (Borisova et al., 2011; see Phosphonoacetate Hydrolase). In addition nine Gram-negative bacterial strains possessed a copy of the phosphonate biosynthetic operon that was flanked by genes encoding enzymes involved in exopolysaccharide biosynthesis; it is known that the capsular polysaccharide B complex associated with virulence in *Bacteroides fragilis* is characterized by its high content of 2-AEP.

**OCCURRENCE OF THE *phnJ* COMPONENT OF THE C–P LYASE OPERON IN SEQUENCED BACTERIAL GENOMES**

Analysis of the distribution of *phnJ*, which encodes a catalytic component of the C–P lyase multi-enzyme system, indicates that C–P lyase is (after “phosphonatase,” see Occurrence in Sequenced Bacterial Genomes of the *phnX* Gene Encoding

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**Table 1 | Polypeptides used as query sequences for bioinformatic analyses.**

| Pathway                        | Enzyme               | Accession number | Microbial origin                | Reference                  |
|--------------------------------|----------------------|------------------|---------------------------------|----------------------------|
| Phosphonate biosynthesis       | Ppd                  | YP_099861.1      | *Bacteroides fragilis* YCH46    | Kuwahara et al. (2004)     |
| C–P lyase                      | Complex subunit PhnJ | NP_418522.1      | *Escherichia coli* str. K-12    | Bury-Mone et al. (2009)    |
| Phosphonoacetalddehyde hydrolase | PhnX                | NP_250002.1      | *Pseudomonas aeruginosa* PAO1   | Cirz et al. (2006)         |
| Phosphonoacetate hydrolase     | PhnA                 | AAC15507.1       | *Pseudomonas fluorescens* 23F   | Kulakova et al. (2001)     |
| Phosphonopyruvate hydrolase    | PalA                 | AAO24736.1       | *Variovorax sp.* Pal2           | Kulakova et al. (2009)     |
| Phosphonate transport          | Transporter subunit PhnD | NP_418529.1   | *Escherichia coli* str. K-12    | Bury-Mone et al. (2009)    |
| Potential new degradation pathway | PhnY*                | ACU83549         | Uncultured bacterium HF130_AEPhn1 | Martinez et al. (2010)   |
| (Martinez et al. (2010))       | PhnZ                 | ACU83550         | Uncultured bacterium HF130_AEPhn1 | Martinez et al. (2010)   |
| Protein of unknown function    | Complex subunit DUF6 | YP_0010234431    | *Methylphilium petroleiphilium* PM1 |                          |
| related to C–P lyase           | Complex subunit DUF1045 | YP_0010234431 | *Acidiphilium cryptum* JF5      |                          |
Table 2 | Homologs retrieved from the NCBI bacterial genomes database that indicate possession of pathways of phosphonate metabolism.

| Pathway                                      | Query polypeptide | “Hits” in sequenced strains | “Hits” in marine strains |
|----------------------------------------------|-------------------|-----------------------------|--------------------------|
| Phosphonate biosynthesis                     | Ppd               | 129 (9.3%)                  | 7 (3.6%)                 |
| C–P lyase                                    | Complex subunit PhnJ | 223 (16.1%)              | 44 (23.2%)              |
| Phosphonoacetaldehyde hydrolase              | PhnX              | 235 (170%)                 | 50 (26.3%)              |
| Phosphonoacetate hydrolase                   | PhnA              | 89 (6.4%)                  | 6 (3.2%)                |
| Phosphonoxyruvate hydrolase                  | PalA              | 3 (0.2%)                   | 0                       |
| Phosphonate transport                        | Transporter subunit PhnD | 294 (21.2%)              | 67 (35.3%)              |
| Potential new 2-AEP degradation pathway      | PhnY*             | 5 (0.4%)                   | 4 (2.1%)                |
|                                               | PhnZ              | 5 (0.4%)                   | 4 (2.1%)                |
| Unknown function related to C–P lyase        | Complex subunit DUF6 | 9 (0.7%)                  | 4 (2.1%)                |
|                                               | Complex subunit DUF1045 | 78 (5.6%)              | 22 (11.6%)              |

*Expressed as a percentage of the 1384 sequenced bacterial genomes contained in the database.

+Expressed as a percentage of the 190 sequenced marine bacterial genomes contained in the database.

Phosphonoacetaldehyde Hydrolase) the second most abundant pathway for phosphonate degradation amongst sequenced bacteria, and that it is widely distributed amongst diverse bacterial groups (Figure 3). Its 223 homologs, including 44 from marine strains, occur mainly in the alpha-, beta-, and gamma-proteobacteria. (Interestingly, however, a copy of the entire operon, and a second copy of its phnCDE genes, were found in the genome of the archaeon Haloquadratum walsbyi; further analysis suggests that these copies have separate proteobacterial origins).

Great diversity was found in the composition and architecture of the C–P lyase operon (including insertions, substitutions, and deletions of components); this confirms a similar observation by Huang et al. (2005) and suggests that the operon has been subject to multiple horizontal gene transfers. All these genetic variations could represent potential variations in the specificity or functionality of the reaction sequence encoded, and thus provide possible evolutionary advantages to their hosts. For example domain of unknown function (DUF) 1045, which is not conventionally designated as one of the 14 genes of the C–P lyase operon, was in fact present in 78 of the 223 operons identified (almost all from the alpha- and beta-proteobacteria, and including 26 of the 44 marine strains in which the operon was found). In 86% of instances DUF1045 flanked or lay close to phnN which is thought to encode a “feeder” activity for the C–P bond cleavage pathway (Hove-Jensen et al., 2011). Expression of both DUF1045 and phnN has been found to be highly induced in the presence of methylphosphonate or the absence of Pi in Acidithiobacillus ferrooxidans (Vera et al., 2008).

Genomic data thus appear to provide support for existing biochemical evidence (see C–P Lyase above) that the C–P lyases may...
be considered as a family of closely related multienzyme systems that nevertheless vary in, for example, their substrate specificities. The high incidence of “optional” genes suggests that these may provide additional functional features. In terms of their regulation, ~90% of C–P lyase operons were apparently controlled by flanking GntR regulators, confirming numerous findings that this pathway is inducible only under conditions of Pi-starvation (Wackett et al., 1987). Surprisingly, however, genes predicted to encode alternative regulators (including those from the LysR, IclR, DeoR, LuxR, MarR, ArsR, TetR, and TonB families) were found adjacent to C–P lyase operons in some 6% of cases, while in a small number of others no putative regulatory genes could be identified.

**OCCURRENCE IN SEQUENCED BACTERIAL GENOMES OF THE phnX GENE ENCODING PHOSPHONOOXALDEHYDE HYDROLASE**

Genes encoding the phosphonatase pathway of 2-AEP degradation, which comprises 2-AEP-pyruvate transaminase (PhnW) and phosphonoacetaldehyde hydrolase (PhnX; see Phosphonoacetaldehyde Hydrolase), were found in 235 bacterial genomes of which 50 were of marine origin, mostly from within the alpha- and gamma-proteobacteria. It is thus the most abundant phosphonate degradation pathway amongst the 1384 bacteria whose sequences are currently available; this may reflect the fact that 2-AEP is thought to be the most widely distributed biogenic phosphonate in the environment (White and Metcalf, 2007). The distribution of genes encoding the phosphonatase operon is shown in Figure 4; some 44% belonged to the gamma-proteobacteria, of whom the majority were potential pathogens (Salmonella, Vibrio, etc.) that have been implicated in gastrointestinal diseases. Predicted phnX-containing strains belonging to the bacillales, bacteroidetes, and clostridia are also associated with similar pathologies. To date, there is no direct evidence to link phosphonoacetaldehyde hydrolase activity to pathogenesis; however it has been reported that low Pi conditions can enhance the virulence of pathogenic species such as P. aeruginosa and Salmonella typhimurium (Beier and Gross, 2006; Lamarche et al., 2008). Additionally, mutations in Pho regulon genes have been thought to be responsible for the decrease of colonization rates by intestinal strains of E. coli and *Vibrio cholerae* (von Krüger et al., 1999, 2006). Since phnX in most gamma-proteobacteria was associated with GntR regulators and hence likely to be expressed under Pi limiting conditions (Gebhard and Cook, 2008), it is conceivable that phosphonoacetaldehyde hydrolase may play a role in the infection process.

The phosphonatase operons in Bacillaceae and clostridial strains were observed to be flanked by deoR, and in bacteroidetes by arac regulatory sequences. Only in alpha-proteobacteria (mostly marine isolates), and in beta-, and gamma-proteobacteria (all pseudomonads) were the phosphonatase pathways associated with LysR type regulators. No such strains have been implicated in gastrointestinal infections. Several features of the phosphonatase operon structures provide strong evidence that PhnX is encoded as part of a genomic island which can be easily transferred amongst bacteria. “Pick-and-mix” horizontal gene transfer is suggested not only by the broad range of regulatory systems apparently involved in the control of the operon expression (including GntR, LysR, DeoR, AraC, and TetR), but also by the insertion of genes such as that encoding 4-aminobutyrate transferase, or DUF6 homologs (the latter is found in some marine strains; DUF6 can also occur within some C–P lyase operons). Furthermore the various rearrangements of the operon components (phnW, X and the phosphonate transporter genes phnC, D, E) that were observed is a feature that is almost entirely absent in pathways of apparently shorter evolutionary history, e.g., that involving phnA (see Occurrence in Sequenced Bacterial Genomes of the phnA Gene Encoding Phosphonoacetic Acid Hydrolase).

**OCCURRENCE IN SEQUENCED BACTERIAL GENOMES OF THE phnA GENE ENCODING PHOSPHONOOXALDEHYDE HYDROLASE**

The phnA gene was much less widely distributed in sequenced bacterial genomes than those encoding the C–P lyase or phosphonoacetaldehyde hydrolase pathways. Figure 5 shows the affiliation of the 69 bacterial strains that encoded the 89 phnA homologs detected during the bioinformatic analysis (these were...
predominantly *Burkholderia* spp. within the beta-proteobacteria). Only six strains were identified as marine isolates; however five of these are members of the *Roseobacter* clade, which is known to be the predominant bacterial group in the near-surface waters of the oceans.

The limited distribution of *phnA* within other phyla suggests that phosphonate degradation via phosphonoacetate hydrolase is a pathway of recent development or that it lacks the transferability observed in the cases of the C–P lyase and phosphonoacetaldehyde hydrolase pathways. Nevertheless, the *phnA* gene was frequently duplicated within the genomes of the species that possessed it (e.g., in 20 different strains of beta-proteobacteria). Only three sequenced strains possessed duplications of *phnX*, and five strains a duplication of the C–P lyase operon. Mechanisms of duplication have been suggested to consolidate the existence of pathways in the host genome, as has been reported for phthalate metabolism in *Rhodococcus* sp. strain DK17 (Choi et al., 2007).

In contrast with the C–P lyase and phosphonatase pathways with their apparently longer evolutionary histories, analysis of those operons that contained *phnA* homologs indicated that these were almost exclusively controlled by only two kinds of regulators: GntR or LysR. Of these, GntR regulators were exclusively encoded in gamma-proteobacteria, with LysR present in all alpha- and some beta-proteobacterial strains (including all six marine strains identified). Rearrangements in the structure of the operons that contained *phnA* were very limited; most (71 out of 89) contained the *phnY* gene that encodes phosphonoacetaldehyde dehydrogenase and *phnW* that encodes 2-AEP aminotransferase, suggesting that phosphonoacetate hydrolase (*phnA*) is more commonly a component of a degradative sequence for the catabolism of 2-AEP (see Phosphonoacetate Hydrolase), than directly inducible by phosphonoacetate as is the case in the operon originally characterized in *P. fluorescens* 23F by Kulakova et al. (2001), in which neither *phnW* or *phnY* is present.

**DISTRIBUTION IN SEQUENCED BACTERIAL STRAINS OF THE** \( pph \) **GENE ENCODING PHOSPHONOPYRUVATE HYDROLASE**

In contrast to the wide distribution exhibited by the phosphonate degradative sequences analyzed above, *pph* was found to be present in only three sequenced bacterial strains, all *Burkholderia* spp.
spp. In each instance the associated operon was controlled by a LysR regulator and contained a copy of the *phnW* gene coding for 2-AEP-aminotransferase. This suggests that the operon associated with phosphonopyruvate hydrolase activity may be of relatively recent origin and that it has not been subject to extensive gene transfer.

**DISTRIBUTION IN SEQUENCED BACTERIAL STRAINS OF ADDITIONAL GENES INVOLVED IN PHOSPHONATE METABOLISM**

In addition to the pathways responsible for the breakdown of phosphonates, the distribution of genes that encode other activities associated with their metabolism was analyzed.

The gene that encodes the PhnD periplasmic binding protein of the phosphonate PhnCDE transporter was very widely distributed across bacteria ([Figure 6](#)), even more so than those encoding the C–P lyase catalytic protein, PhnJ, or phosphonoacetaldehyde hydrolase. Numbers of predicted homologs were particularly high amongst marine bacteria; of the 294 strains in the NCBI database that were confirmed to possess a *phnD* homolog, 67 were of marine origin. In 177 of the strains identified, *phnD* was juxtaposed with a phosphonate degradative gene (thus 60% of strains capable of transporting phosphonates into the cell have an apparent ability to degrade them). On the other hand, 117 other strains that contained *phnD* did not encode any known pathway of phosphonate degradation, suggesting that not all phosphonate degradation pathways have yet been discovered (as has been proposed by Palenik et al., 2003). It is also possible that phosphonates are not always used as a P source, but after transport into the cell may be incorporated into phospholipids, phosphonoproteins, or other macromolecules by anabolic routes. And conversely, 289 strains that possessed genes associated with a known phosphonate degradation pathway showed no evidence of possessing the PhnCDE route of phosphonate acquisition – suggesting the existence of as-yet unknown uptake routes.

An apparent example of a recent pathway of phosphonate metabolism is encoded by the operon designated as PhnY*/PhnZ* and discovered by Martinez et al. (2010) in a clone from a fosmid library from the Sargasso Sea. The enzymes of the pathway have not yet been characterized, but it was experimentally proven that they conferred the capacity to consume phosphonates as a P source on an *E. coli* *phn* mutant strain (Martinez et al., 2010). PhnY* is thought to be a 2-oxoglutarate dioxygenase, and PhnZ a protein of the HD phosphohydrolase family (Martinez et al., 2010). Bioinformatic analysis indicated that only five sequenced bacterial strains possessed a copy of this operon. Interestingly, four of these were isolated from the marine environment, suggesting that this pathway may have evolved in the oceans and may now play a significant role in P acquisition in marine microorganisms. Three of the strains (*Pleisoicytis pacifica* SIR-1, and *Prochlorococcus marinus* str. MIT9301 and MIT9303) possess a copy of the phosphonate transporter operon *phnCDE* but had been considered to lack any recognized phosphonate degradation pathway. As is the case with the phosphonopyruvate hydrolase pathway of C–P bond cleavage, the PhnY*/PhnZ* pathway appears to have undergone no extensive evolutionary rearrangement. In this paper the *phnY* gene described by Martinez et al. (2010) is designated as *phnY*.

**ABUNDANCE AND DISTRIBUTION OF THE GENES OF PHOSPHONATE METABOLISM IN MARINE METAGENOMIC DATABASES**

The analysis presented in Section “Distribution of the Genes of Phosphonate Metabolism Amongst Sequenced Bacteria” is based on the information currently available on sequenced bacterial genomes which remains heavily biased in favor of species of clinical, or other applied significance. Hence little can be extrapolated to the ecological significance of the various pathways of phosphonate metabolism identified. In an attempt to address this shortcoming, query sequences indicative of the various phosphonate metabolic pathways were used to interrogate the specialist genomic database for marine bacteria, CAMERA (Nealson and Venter, 2007), using the protocol of Lim et al. (2007). An indication was thus obtained of the abundance of the genes encoding those pathways in genomic DNA recovered at the 83 sites from the surface of the world’s oceans sampled by the global

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**FIGURE 6** | Distribution of *phnD* homologs within sequenced bacterial genomes. The numbers and affiliations of marine bacterial strains that contain the *phnD* gene, out of the 190 sequenced marine bacterial genomes publically available, are indicated to the right of the Figure.
ocean survey (GOS) Expedition (Table 3). The results from the CAMERA BLAST analysis (including those for some additional activities associated with bacterial P metabolism) were also normalized against the number of “hits” obtained for RecA; *recA* is frequently used as a single-copy-per-genome control in distribution and prevalence analyses (Moran et al., 2004; Karl et al., 2008; Table 4).

The *ppd* gene that encodes the phosphonate biosynthetic enzyme phosphonopyruvate decarboxylase was found at 67 sites, almost 81% of those sampled by the GOS Expedition, and in the equivalent of 10% of sampled genomes. This suggests that natural-product phosphonates may be widely produced in the oceanic environment, and correlates with the potential biological advantages that these molecules may provide as structural components or secondary metabolites. To date, no experimental data has been presented on the ability of marine bacterial strains to synthesize phosphonates, although a recent study using solid-state NMR spectroscopy has demonstrated that C–P compounds comprised almost 10% of the cellular particulate phosphorus pool in cultured strains of *T. erythraeum*, suggesting that this cyanobacterium might be capable of C–P bond formation (Dyhrman et al., 2009).

In respect of phosphonate catabolism, bioinformatic analysis indicated a wide distribution of the phosphonate transporter subunit, PhnD. The confirmed presence of the *phnD* gene at 72% of the sites, and in some 5% of the genome-equivalents predicted, suggests that phosphonates are not only being synthesized by bacteria in the oceans but also that exogenous C–P compounds can be taken up for catabolism (or other metabolic activities) at significant levels. This is consistent with recent *in situ* expression studies on *phnD* in marine picocyanobacteria by Ilikchyan et al. (2009).

In total, some 28% of bacterial genome-equivalents sampled during the GOS were found to contain genes encoding one or more pathways of C–P bond cleavage (Table 4); however the apparent relative abundances of phosphonate degradation pathways contrasted with those found amongst sequenced bacteria in the GenBank database. Thus genes encoding C–P lyase and phosphonoacetaldehyde hydratase had a more limited presence in the oceanic environment, being present at only 30 and 29% respectively of the sites sampled by the GOS (Table 3), and in 5 and 2% of sampled genomes (Table 4). However the *phnA* gene that encodes phosphonoacetate hydratase was more widely distributed in the oceans, being present at almost over 78% of the sites sampled by the GOS Expedition (Table 3), and in an estimated 11% of genome-equivalents (Table 4). Similarly, the genes that encode the PhnY*/PhnZ phosphonate catabolic pathway (Martinez et al., 2010) were of limited occurrence in sequenced bacterial genomes but were amongst the most widely distributed of those analyzed in the GOS database; as noted above, four of the five

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**Table 3 | Presence of genes encoding pathways of phosphonate metabolism at the 83 oceanic sites sampled by the GOS Expedition.**

| Pathway                                | Query polypeptide                  | % of sampled GOS sites |
|----------------------------------------|------------------------------------|------------------------|
| Phosphonate biosynthesis               | Phosphonopyruvate decarboxylase    | 80.7                   |
| C–P lyase                              | Complex subunit PhnJ               | 30.1                   |
| Phosphonoacetaldehyde hydratase        | PhnX                               | 28.9                   |
| Phosphonoacetate hydratase             | PhnA                               | 78.3                   |
| Phosphonopyruvate hydratase            | PalA                               | 1.2                    |
| Phosphonate transporter                | Transporter subunit PhnD           | 72.3                   |
| Potential new degradation pathway      | PhnY*                              | 51.8                   |
|                                       | PhnZ                               | 77.1                   |
| Repair and maintenance of DNA (Control)| RecA                               | 100                    |

**Table 4 | Abundance of phosphonate (and related) metabolic pathways in bacterial genomes sampled by the Global Ocean Survey.**

| Pathway                                | Query polypeptide                  | Abundance of pathway (%) |
|----------------------------------------|------------------------------------|--------------------------|
| Phosphonate biosynthesis               | Phosphonopyruvate decarboxylase    | 10.2                     |
| C–P lyase                              | Complex subunit PhnJ               | 5.1                      |
| Phosphonoacetaldehyde hydratase        | PhnX                               | 1.8                      |
| Phosphonoacetate hydratase             | PhnA                               | 11.2                     |
| Phosphonopyruvate hydratase            | PalA                               | 0.1                      |
| Phosphonate transporter                | Transporter subunit PhnD           | 5.5                      |
| Potential new degradation pathway      | PhnY*                              | 1.6                      |
|                                       | PhnZ                               | 9.4                      |
| Alkaline phosphatase A                 | PhoA                               | 5.9                      |
| Alkaline phosphatase D                 | PhoD                               | 4.6                      |
| Phosphatase X                          | PhoX                               | 3.5                      |
| Repair and maintenance of DNA (Control)| RecA                               | 100                      |
sequenced isolates that possess \textit{phnY}*/\textit{phnZ} are marine strains (see Distribution in Sequenced Bacterial Strains of Additional Genes Involved in Phosphonate Metabolism).

This discrepancy between the apparent abundances of the various pathways of phosphonate catabolism and their prevalence in sequenced bacterial genomes may be in part explained by the fact that the majority of sequenced strains that possess a copy of the C–P lyase and phosphonoacetaldehyde hydrolase operons belong to the gamma-proteobacteria and other classes of prokaryotes associated with gastrointestinal pathogenesis, whereas those strains found to possess the phosphonoacetate hydrolase and PhnY*/PhnZ pathways were in many cases alpha-proteobacteria. Most of these had been isolated from the marine environment; they comprise the predominant class of prokaryotes in the upper layer of the sea surface (Wagner-Döbler and Biebl, 2006; Moran et al., 2007). Furthermore, since all those sequenced marine strains that possessed a \textit{phnA} (or a \textit{phnX}) – encoded pathway of 2-AEP degradation had an associated \textit{lyS}R gene encoding a transcriptional activator it is possible that this form of inducible regulation might be of competitive advantage, allowing cells to consume phosphonates as an additional source of C, (N), and P independently of ambient Pi concentrations. Given that expression of the genes that encode the C–P lyase operon in all cases studied appears to Pi-starvation-inducible, possession of a Pi-insensitive mechanism of phosphonate catabolism could be of particular relevance in niches of localized organic enrichment within the bulk oligotrophic oceans, such as “marine snow.”

The limited available data from other studies are broadly consistent with our findings. Thus a metagenomic study of a coastal marine mesocosm by Gilbert et al. (2009) found that \textit{phnA} was contained in >3% of genomes sampled while analysis of a metagenomic database derived from a 4000-m depth profile sampled in the North Pacific Subtropical Gyre (Luo et al., 2011) found that \textit{phnA} was present in approximately 32% of genomes sampled throughout the water column, but that C–P lyase genes were confined to P-limited surface waters. Martinez et al. (2010) in metagenomic analyses of depth profile samples from the Sargasso Sea and Hawaii Ocean Time Series found that \textit{phnA} was particularly abundant in deeper waters (present in an average of 10% of genomes across all depths at the two sites). The \textit{phnY}* and \textit{phnZ} genes were present on average in 3 and 10% respectively of genomes at these sites (Martinez et al., 2010). Finally, the likely relative importances of phosphonate-P and ester-P within marine DOP can perhaps be gauged by comparison of the broadly similar abundances in GOS metagenomic samples of genes that encode C–P cleavage enzymes and those that encode the three major marine alkaline phosphatases (Table 4).

**SUMMARY/CONCLUSION**

There is growing awareness of the importance of phosphonates in the P economy of the oceans, although analytical constraints continue to limit our knowledge of the identities, abundances, and biogenic origins of marine C–P compounds, and of the dynamics of their biotransformations. Genomic, metagenomic (and increasingly, metatranscriptomic) evidence all point to the widespread utilization of phosphonate-P by the marine microflora; the indications these provide that regulation of C–P bond cleavage activity may be niche-dependent require further exploration. In addition there is need for further experimental proof of strong in silico suggestions that marine bacteria may be of importance as C–P producers, in addition to their better-known role as phosphonate consumers.

**ACKNOWLEDGMENTS**

We thank CONACYT, Mexico, for a studentship to Juan F. Villarreal-Chiu.

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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.

Received: 30 November 2011; paper pending published: 22 December 2011; accepted: 10 January 2012; published online: 26 January 2012.

Citation: Villarreal-Chiu IF, Quinn JP and McGrath JW (2012) The genes and enzymes of phosphonate metabolism by bacteria, and their distribution in the marine environment. Front. Microbiol. 3:19. doi: 10.3389/fmicb.2012.00019

This article was submitted to Frontiers in Aquatic Microbiology, a specialty of Frontiers in Microbiology.

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