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#### Authors

Youssef, Noha H

Rinke, Christian

Stepanauskas, Ramunas

et al.

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## ORIGINAL ARTICLE

# Insights into the metabolism, lifestyle and putative evolutionary history of the novel archaeal phylum ‘Diapherotrites’

Noha H Youssef<sup>1</sup>, Christian Rinke<sup>2</sup>, Ramunas Stepanauskas<sup>3</sup>, Ibrahim Farag<sup>1</sup>, Tanja Woyke<sup>2</sup> and Mostafa S Elshahed<sup>1</sup>

<sup>1</sup>Department of Microbiology and Molecular Genetics, Oklahoma State University, Stillwater, OK, USA;

<sup>2</sup>DOE Joint Genome Institute, Walnut Creek, CA, USA and <sup>3</sup>Bigelow Laboratory for Ocean Sciences, East Boothbay, ME, USA

The archaeal phylum ‘Diapherotrites’ was recently proposed based on phylogenomic analysis of genomes recovered from an underground water seep in an abandoned gold mine (Homestake mine in Lead, SD, USA). Here we present a detailed analysis of the metabolic capabilities and genomic features of three single amplified genomes (SAGs) belonging to the ‘Diapherotrites’. The most complete of the SAGs, *Candidatus ‘Iainarchaeum andersonii’* (Cand. IA), had a small genome (~1.24 Mb), short average gene length (822 bp), one ribosomal RNA operon, high coding density (~90.4%), high percentage of overlapping genes (27.6%) and low incidence of gene duplication (2.16%). Cand. IA genome possesses limited catabolic capacities that, nevertheless, could theoretically support a free-living lifestyle by channeling a narrow range of substrates such as ribose, polyhydroxybutyrate and several amino acids to acetyl-coenzyme A. On the other hand, Cand. IA possesses relatively well-developed anabolic capabilities, although it remains auxotrophic for several amino acids and cofactors. Phylogenetic analysis suggests that the majority of Cand. IA anabolic genes were acquired from bacterial donors via horizontal gene transfer. We thus propose that members of the ‘Diapherotrites’ have evolved from an obligate symbiotic ancestor by acquiring anabolic genes from bacteria that enabled independent biosynthesis of biological molecules previously acquired from symbiotic hosts. ‘Diapherotrites’ 16S rRNA genes exhibit multiple mismatches with the majority of archaeal 16S rRNA primers, a fact that could be responsible for their observed rarity in amplicon-generated data sets. The limited substrate range, complex growth requirements and slow growth rate predicted could be responsible for its refraction to isolation.

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## Introduction

Our view of the phylogenetic diversity and ecological distribution of members of the domain Archaea is rapidly evolving. Historically, Archaea were regarded as a collection of extremophiles that could only thrive in extreme habitats, for example, high temperature (Huber *et al.*, 1991; Jannasch *et al.*, 1992; Antoine *et al.*, 1995; Whitaker *et al.*, 2003), high salinity (Oren *et al.*, 1990; Cui *et al.*, 2010; Goh *et al.*, 2011; Inoue *et al.*, 2011), low pH (Edwards *et al.*, 2000; Dopson *et al.*, 2004), strict anaerobic conditions (Mikucki *et al.*, 2003; Sakai *et al.*, 2007) or combinations thereof (Huber *et al.*, 1989; Itoh *et al.*, 1999; Minegishi *et al.*, 2008, 2013).

This notion was subsequently challenged by culture-independent diversity surveys that documented the occurrence of the Archaea in both extreme (Bond *et al.*, 2000; Orphan *et al.*, 2000; Benlloch *et al.*, 2001, 2002; Baker and Banfield, 2003) and temperate habitats, for example, soil (Bintrim *et al.*, 1997; Walsh *et al.*, 2005; Bates *et al.*, 2011; Tripathi *et al.*, 2013), marine environments (DeLong, 1992; Fuhrman *et al.*, 1992; Massana *et al.*, 1997; Karner *et al.*, 2001) and freshwater ecosystems (Lin *et al.*, 2012; Yergeau *et al.*, 2012; Berdjeb *et al.*, 2013; Bricheux *et al.*, 2013; Silveira *et al.*, 2013; Vila-Costa *et al.*, 2013). In addition to establishing the ubiquity of the Archaea on a global scale, these studies have also significantly expanded the scope of phylogenetic diversity within this domain, as many of the sequences identified represented novel, high-rank phylogenetic lineages (DeLong, 1992; Vetriani *et al.*, 1999; Takai *et al.*, 2001; Hallam *et al.*, 2004; Elkins *et al.*, 2008; Hu *et al.*, 2011; Nunoura *et al.*, 2011).

Correspondence: NH Youssef, Department of Microbiology and Molecular Genetics, Oklahoma State University, 1110S Innovation Way Drive, Stillwater, OK 74074, USA.

E-mail: noha@okstate.edu

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Environmental genomic approaches and dedicated isolation efforts have yielded valuable insights into the metabolic capabilities and ecological roles of many of these novel lineages (Hallam *et al.*, 2004; Konneke *et al.*, 2005; Elkins *et al.*, 2008; Walker *et al.*, 2010; Lloyd *et al.*, 2013). Recently, the pace of discovery and characterization of archaeal lineages has significantly accelerated, driven by recent methodological and computational advances in single cell sorting and amplification, sequencing methodologies and novel binning and assembly approaches that enable efficient genomic reconstruction from metagenomic sequence data (Dick *et al.*, 2009; Wrighton *et al.*, 2012; Rinke *et al.*, 2013; Swan *et al.*, 2013). These advances led to the identification and genomic characterization of multiple novel high-rank archaeal lineages that have previously escaped detection in 16S ribosomal RNA (rRNA) gene-based diversity surveys because of their extremely low relative abundance, limited distribution or mismatches to archaeal 16S rRNA gene primer sequences (Baker *et al.*, 2010; Nunoura *et al.*, 2011; Narasingarao *et al.*, 2012). These discoveries necessitated phylogenetic and phylogenomic-based reassessment of the taxonomic structure of the domain *Archaea* (Lake *et al.*, 1984; Brochier-Armanet *et al.*, 2008; Elkins *et al.*, 2008; Ghai *et al.*, 2011; Guy and Ettema, 2011; Williams *et al.*, 2012). The most recent and comprehensive phylogenomics-based assessment of the domain *Archaea* (Rinke *et al.*, 2013) combined data from 35 novel archaeal single amplified genomes (SAGs) with previously published archaeal genomes to propose a three archaeal superphyla scheme. These superphyla are the Euryarchaeota, the TACK superphylum (encompassing the Thaumarchaeota, 'Aigarchaeota', Crenarchaeota and Korarchaeota as previously suggested; Guy and Ettema, 2011; Williams *et al.*, 2012) and the newly proposed DPANN superphylum.

The DPANN superphylum encompasses the 'Nanoarchaeota' (Waters *et al.*, 2003; Podar *et al.*, 2013), the only DPANN phylum with cultured representatives, as well as the candidate phyla 'Nanohaloarchaeota' (defined from metagenomic assembly (Narasingarao *et al.*, 2012) and SAGs (Ghai *et al.*, 2011) from hypersaline environments); 'Parvarchaeota' (defined from a metagenomic assembly from an acid mine drainage; Baker *et al.*, 2010), 'Aenigmarchaeota' (defined from three SAGs from Homestake mine groundwater seep (Lead, SD, USA) and the Great Boiling Spring sediments; Rinke *et al.*, 2013) and 'Diapherotrites' (defined from SAGs from Homestake mine groundwater seep; Rinke *et al.*, 2013). As such, the DPANN superphylum represents an intriguing collection of phyla with disparate physiological preferences and environmental distribution, ranging from the obligatory symbiotic and thermophilic species within the 'Nanoarchaeota', to the acidophilic candidate phylum 'Parvarchaeota' and to the non-extremophilic candidate phyla 'Aenigmarchaeota' and 'Diapherotrites'.

Here, we present a detailed analysis of the genomic, metabolic and ecological features of three SAGs belonging to the 'Diapherotrites'. We present evidence of genomic streamlining as well as limited metabolic capacities within the analyzed genomes. We also demonstrate the prevalence of cross-kingdom horizontal gene transfer (HGT) events, and argue that HGT process represents an important evolutionary mechanism that contributes to the observed genomic features, ecological distribution and proposed trophic lifestyle of members of this phylum.

## Materials and methods

### *Origin of Diapherotrites SAGs*

Candidate phylum 'Diapherotrites' (CP-'Diapherotrites') SAGs analyzed in this study were all obtained from a groundwater seep from the ceiling of Homestake Mine (Lead, SD, USA) drift at a depth of 100 m as described previously (Rinke *et al.*, 2013). Cell sorting and lysis, single-cell whole genome amplification, 16S rRNA amplicon sequencing as well as SAG sequencing and assemblies have been detailed before (Rinke *et al.*, 2013). Three CP-'Diapherotrites' SAGs were deposited under Genbank assembly IDs GCA\_000402355.1, GCA\_000404545.1 and GCA\_000404525.1, and Integrated Microbial Genomes (IMG) SAG names SCGC\_AAA011-E11, SCGC\_AAA011-K09 and SCGC\_AAA011-N19. The candidatus-type species for CP-'Diapherotrites' is the SAG SCGC\_AAA011-E11, for which the name *Candidatus* 'Iainarchaeum andersonii' was proposed (Rinke *et al.*, 2013). Our analysis was mainly conducted on the *Candidatus* Iainarchaeum andersonii genome (henceforth referred to as Cand. IA), as it had the highest estimated genome completion (88.5%) among the three CP-'Diapherotrites' genomes (Supplementary Table S1), with the other two partial SAGs mainly used for confirmatory purposes.

### *Genome annotation, general genomic features and metabolic reconstruction*

Genome functional annotation was performed using the IMG platform (<http://img.jgi.doe.gov>) as previously described (Rinke *et al.*, 2013). Metabolic reconstruction was conducted using both KEGG (Kyoto Encyclopedia of Genes and Genomes) and Metacyc databases (Kanehisa, 2002; Karp *et al.*, 2002). Proteases, peptidases and protease inhibitors were predicted using Blastp against the Merops database (Rawlings *et al.*, 2014). Transporters were identified by querying the genome against the TCDB (transporter classification database) (Saier *et al.*, 2014) using Blastp (Altschul *et al.*, 1990). Gene duplications were identified by running local Blastp using the proteins as both the subject and the query, where non-self hits with a similarity cutoff of >90% are considered duplicates. Overlapping genes were

identified by comparing the coordinates of the start and stop codons for all genes on the same contig. Protein subcellular localizations were identified online using the PSORTb V 3.0.2 (Yu *et al.*, 2010). Protein COG (Cluster of orthologous groups; Tatusov *et al.*, 2000) family distributions for Cand. IA and reference genomes were either obtained from the corresponding IMG genome page or, in case the genome was not available in the IMG database, were identified using the web batch conserved domain (Marchler-Bauer *et al.*, 2010) search tool (available at <http://www.ncbi.nlm.nih.gov/Structure/bwrpsb/bwrpsb.cgi>) against the COG database with an *E*-value threshold of 0.001.

#### *HGT in Cand. IA genome assembly*

A two-tier approach was utilized to evaluate the putative frequency of occurrence of HGT in Cand. IA genome assembly. Blastp (Altschul *et al.*, 1990) was conducted for all genes in Cand. IA as well as other representatives of the DPANN superphylum. Genes were classified according to the Blastp first hit. For in-depth evaluation of HGT within Cand. IA, 21 HGT candidate genes with bacterial Blastp first hit were chosen for further analysis according to the following two criteria: (1) protein length is longer than 100 amino acids and (2) placement on a contig with other genes involved in the same pathway but of apparent archaeal origin. In cases where all genes involved in a pathway had bacterial Blastp first hit, priority was for genes present on longer contigs with other genes of apparent archaeal origin. Homologs of genes of interest were obtained from Genbank by BLASTP of the target gene. The obtained homolog data sets were comparatively aligned using COBALT (Papadopoulos and Agarwala, 2007). A guide tree was used to select the closest relatives of the target gene. In cases where all the closest relatives belonged to bacterial phyla, archaeal counterparts were obtained from Genbank and included in the downstream analysis. A eukaryotic outgroup was also included. For phylogenetic tree construction, target genes and their homologs were aligned using ClustalW (Larkin *et al.*, 2007). Maximum likelihood trees were calculated from the alignment using the BLOSUM62 model and GAMMA approximation implemented in RaxML (Stamatakis, 2014).

#### *Principal component analysis (PCA)*

PCA was conducted to identify salient differences in genomic features and COG distributions between DPANN and other model archaeal genomes. A list of genomes, as well as features used in this analysis, is presented in Supplementary Table S2. Initially, all genomic features previously implied as determinants of genome streamlining (Giovannoni *et al.*, 2005; Lauro *et al.*, 2009; Swan *et al.*, 2013) were included in the analysis. These include: genome size, number of genes, GC content, noncoding density, number of ribosomal RNA operons, percentage of

transporter proteins per genome size, COG categories distribution (that is, percentage of proteins belonging to each COG category), protein sublocalization (percentage of proteins destined to the cytoplasm, cytoplasmic membrane, cell wall and extracellular milieu) and encoded amino acid frequency. Data were then implemented in the PCA using the *prcomp* function in the *labdsv* package (Roberts, 2012) of R (R Development Core Team, 2008). A biplot was constructed using the *biplot* function in R, where genomes are represented as points and variables are represented as arrows pointing in the direction of maximal abundance (R Development Core Team, 2008). To simplify the biplots, all variables that showed minimal effect on the PCA biplots (for example, variables whose arrows clustered at the origin) were removed and the analysis was repeated.

*Mismatches in CP-‘Diapherotrites’ 16S rRNA genes to archaeal primers and secondary structure prediction*  
16S rRNA genes from the three CP-‘Diapherotrites’ SAGs were aligned to reference small subunit rRNA sequences, and sites of universal archaeal primers previously utilized in culture-independent surveys were examined to identify putative mismatches and/or indel occurrences. Secondary structure prediction was achieved using the Mfold web server (Zuker, 2003), with the minimum energy structure predicted compared with the conserved secondary structure of both the *Escherichia coli* and *Methanospirillum hungatei* 16S rRNA molecules.

#### *Mining public databases to elucidate the distribution of the ‘Diapherotrites’ in nature*

We attempted to identify the occurrence of members of the CP-‘Diapherotrites’ in various ecosystems by mining metagenomic data sets in the IMG database ( $n=893$ , accessed in December 2013), Sanger-generated 16S rRNA gene sequences in the nr database ( $n=53\,650\,62$  sequences, accessed in January 2014) and partial, high-throughput (pyrosequencing and Illumina)-generated archaeal 16S rRNA gene sequences in MG-RAST (Meyer *et al.*, 2008) and SRA archive (Leinonen *et al.*, 2011) ( $n=31\,972\,882$  sequences in 775 data sets generated using archaeal primers). Identification of CP-‘Diapherotrites’ in metagenomic data sets was conducted using the three ‘Diapherotrites’ SAG assemblies for anchoring metagenomic reads as previously described (Rinke *et al.*, 2013). To identify the distribution using data in Sanger- and high-throughput-generated data sets, we followed the protocols previously described in Farag *et al.* (2014).

## Results

### *General features of ‘Diapherotrites’ genomes*

The general features of Cand. IA partial genome compared with sequenced representatives of the

**Table 1** Genomic features of *Candidatus* 'Iainarchaeum andersonii' genome compared with other DPANN superphylum members

Phylum Genome	<i>Diapherotrites</i> <i>Candidatus</i> 'Iainarchaeum andersonii'	<i>Nanoarchaeota</i> <i>Nanoarchaeum</i> <i>equitans</i> <i>Kin-4</i>	<i>Parvarchaeota</i> <i>Candidatus</i> ' <i>Micrarchaeum</i> <i>acidiphilum</i> ' <i>ARMAN-2</i>	<i>Nanohaloarchaeota</i> <i>Candidatus</i> ' <i>Nanosalina</i> ' sp. <i>J07AB43</i>	<i>Aenigmarchaeota</i> <i>Candidatus</i> ' <i>Aenigmarchaeum</i> <i>subterraneum</i> '
Genome size, Mb	1.1	0.49	~1	1.23	0.43
Genome completeness (%)	88.5	100	~99	~100	47.3
Estimated size (Mb)	1.24	0.49	1	1.23	0.91
GC %	47	31.6	47.2	44	38.6
% Noncoding	9.56	7.6	9.3	12.4	5.24
% Overlapping genes	27.63	44.1	18-19	24.9	19.42
Gene duplication* (%)	2.16	ND	ND	ND	ND
Average gene length, bp	822	827	892	639	766
<i>RNA genes</i>					
5S rRNA count	0	1	0	1	2
16S rRNA count	1	1	1	1	1
23S rRNA count	1	1	1	0	0
tRNA count	39	38	35	59	36
Number of CDS	1202	556	1033	1678	525
With function prediction	782	549	668	773	308
Without function prediction	420	7	365	905	217
Number of CDS with inteins	5	1	1	1	0
Number of CDS with IS elements	24	0	10	9	3
Number of paralog clusters	156	53	ND	198	220
<i>% Of proteins sublocalized to</i>					
Cytoplasm	79.9	63.3	65.6	68.9	61.5
Cytoplasmic membrane	14.6	12.3	18.9	12.8	19.23
Extracellular milieu	1.4	0.8	1.2	2.3	1.7

Abbreviations: CDS, coding sequence; IS elements, insertion sequence elements; ND, not determined; rRNA, ribosomal RNA; tRNA, transfer RNA.  
\*Percentage of genes having duplicates with protein sequence similarity  $\geq 90\%$ .

DPANN superphylum are shown in Table 1. *Cand. IA* exhibits general genomic features previously observed in DPANN genomes (Baker and Banfield, 2003; Waters *et al.*, 2003; Narasingarao *et al.*, 2012; Rinke *et al.*, 2013). It has a relatively small genome (estimated size ~1.24 Mb), short average gene length (822 bp), single ribosomal RNA operon, high coding density (~90.4%), high percentage of overlapping genes (27.6%, mean overlap 4 bp, range 1–12 bp) and very low incidence of gene duplication (2.16%).

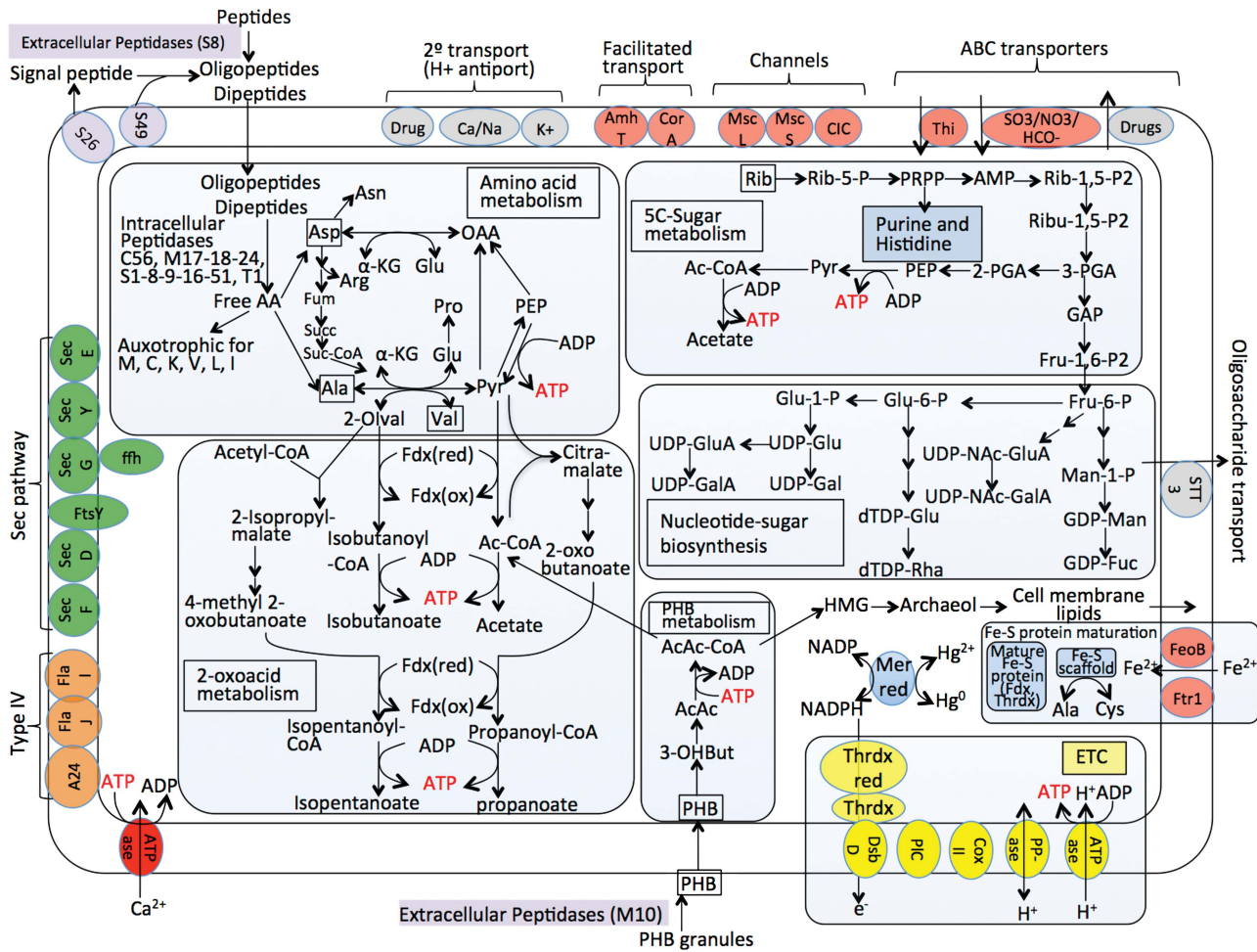
#### Metabolic features of *Cand. IA*

**Catabolic capacities.** Analysis of *Cand. IA* genome suggests that it possesses a relatively limited catabolic potential. The genome lacks evidence for a complete glycolysis, tricarboxylic acid cycle or short-chain fatty acid or alcohol degradation. Nevertheless, the genome suggests the capability to channel few distinct substrates (valine, alanine, aspartate, ribose and polyhydroxybutyrate) into acetyl- (or acyl-) coenzyme A (CoA), and subsequently generate adenosine triphosphate (ATP) via acetyl-CoA synthase (Figure 1). Details of the pathways involved in these processes are provided in the Supplementary Text. In brief, for amino acid degradation, the genome harbors multiple

transaminases that allow for the conversion of alanine, valine and aspartate to the corresponding 2-oxoacids. Oxidative decarboxylation of the 2-oxoacids to acyl-CoA can occur via pyruvate/2-oxoacid:ferredoxin oxidoreductases, or oxaloacetate-decarboxylating malate dehydrogenase (EC: 1.1.1.38) followed by conversion into the corresponding free acid via acyl-CoA synthetase with the concurrent production of 1 ATP. An incomplete tricarboxylic acid cycle could potentially replenish 2-ketoglutarate for the transamination reactions.

*Cand. IA* also appears to possess the capability for ribose degradation (Figure 1), where ribose could potentially be activated by ribokinase and ribose-5-phosphate pyrophosphokinase. The presence of type III ribulose-1,5-bisphosphate carboxylase (Rubisco) in *Cand. IA* genome suggests the employment of the novel adenosine monophosphate (AMP) metabolism pathway suggested by Aono *et al.* (2012). The combination of ribose activation and AMP salvage enzymes eventually lead to the production of 3-phosphoglycerate that feeds into the lower arm of glycolysis and subsequently leads to ATP production as described above. The process results in a net production of 2 ATP/ribose (Supplementary Text).

Another potential carbon and energy source for *Cand. IA* is polyhydroxybutyrate (PHB), the storage compounds produced by many bacteria under



**Figure 1** Metabolic reconstruction for *Candidatus Iainarchaeum andersonii* genome. Double lines surrounding the cell depict cell membrane. Possible catabolic ATP-producing pathways are shown in light blue boxes, and sites of ATP production are shown in red. Potential substrates are shown in boxes, and include the amino acids alanine, valine and aspartate, the 5C-sugar ribose and polyhydroxybutyrate. AcAc, acetoacetate; AcAc-CoA, acetoacetyl-CoA; Fdx, ferredoxin; 3-OHBut, 3-hydroxybutyrate. Electron transport chain components are shown in yellow. ATPase, V-type ATPase pump; CoxII, cytochrome oxidase subunit II; DsbD, disulfide bond oxidoreductase D; PIC, plastocyanin; PPase, inorganic pyrophosphatase; Thrdx, thioredoxin; Thrdx red, thioredoxin reductase. NADPH could potentially act as the electron donor to the ETC. Possible sites of NADPH production in the cell include mercuric reductase enzyme (Mer Red). All predicted transporters with known functions are shown. (1) Secretory pathways: components of the Sec pathway are shown in green, whereas components of type IV pili assembly are shown in orange. (2) Transporters: exporters are shown in gray including secondary antiporters, ABC transporters, STT3 the oligosaccharide exporter, whereas importers are shown in red including the facilitated transporters AmhT and CorA, the channels of MscS, MscL and CIC families, ABC transport systems for thiamine and sulfonate/nitrate/bicarbonate as well as ferrous iron transporters FeoB and Ftr1 for Fe-S assembly. AmhT, for ammonium; CIC, chloride channel families; CorA, for cobalt; MscL, large mechanosensitive; MscS, small mechanosensitive. Peptidases are shown in purple.

conditions of excess carbon. The genome encodes a PHB depolymerase downstream of, and overlapping with, a transmembrane matrixin-coding gene (peptidase family M10) (Visse and Nagase, 2003; Rawlings *et al.*, 2014), known to hydrolyze extracellular matrix components. Therefore, it appears that matrixin is used by *Cand. IA* to break down the protein shell of PHB granules, and the released PHB is subsequently depolymerized into  $\beta$ -hydroxybutyrate. The produced  $\beta$ -hydroxybutyrate could potentially be oxidized to acetoacetate (using a NAD-dependent dehydrogenase belonging to the GFO/IDH/MOCA family (pfam 01408)), followed by activation of acetoacetate to acetoacetyl-CoA using acyl-CoA synthase. The concerted action of

acetyl-CoA C-acetyltransferase and acetyl-CoA synthetase converts acetoacetyl-CoA to two molecules of acetate with the concomitant production of ATP.

Finally, it is plausible that *Cand. IA* might also employ a modified electron transport chain, similar to that recently suggested for candidate division TM6 (McLean *et al.*, 2013). The chain will involve disulfide bond oxidoreductase D (DsbD), a thioredoxin-disulfide (NADPH) reductase (EC. 1.8.1.9) and thioredoxins as initial substrate (NADPH) oxidoreductases, plastocyanins as potential cytochrome equivalents and cytochrome *c* oxidase as a potential terminal oxidase. An inorganic pyrophosphatase and all subunits of V-type ATP synthase

could potentially employ the proton motive force generated across the membrane for ATP synthesis (Figure 1 and Supplementary Text).

**Anabolic potential.** Although the catabolic potential of Cand. IA appears to be rather limited, the genome suggests a fairly well-developed anabolic machinery. Cand. IA genome suggests the capacity for biosynthesis of multiple carbohydrates, amino acids, lipids, nucleotides as well as several cofactors (Figure 1, Table 2 and Supplementary Table S3). Cand. IA genome encodes a partial gluconeogenic pathway up to the level of fructose-6-phosphate. The genome shows evidence for synthesizing most amino acids with the exception of lysine, cysteine, methionine and branched chain amino acids (Figure 1 and Table 2). In addition, all enzymes essential for archaeol, the archaeal membrane lipid, biosynthesis from acetyl-CoA via the mevalonate pathway are encoded in the genome. Cand. IA genome shows evidence for biosynthesis of some cofactors (riboflavin, pyridoxal phosphate, nicotinate and nicotinamide, coenzyme A and ferredoxin) and nucleotides (complete evidence for *de novo* biosynthesis of pyrimidine nucleotides, partial evidence for purines). Finally, all necessary enzymes for assembly of Fe-S and maturation of Fe-S proteins are also encoded by the genome.

**Transporters and extracellular peptidases.** Although the number of transporters encoded by Cand. IA genome are fairly limited ( $n=54$ ), they appear to be able to uptake several cations and anions (Supplementary Text). Several transporters belonging to transporter families with unidentified substrates are annotated as membrane proteins with unknown function and hence might be involved in the transport of amino acids, sugars or cofactors for which Cand. IA appears to be auxotrophic. The genome also encodes for various extracellular and membrane-associated peptidases (see Supplementary Text for more details) that presumably act to cleave extracellular peptides, with the resulting oligopeptides and amino acids subsequently transported into the cell and used to supplement auxotrophies and/or ATP production.

#### *HGT in Cand. IA*

The metabolic analysis described above clearly indicates that Cand. IA genome possesses relatively limited metabolic capabilities as compared with free-living archaeal copiotrophs and oligotrophs. Nevertheless, unlike the model obligate archaeal symbionts within the 'Nanoarchaeota' ('*Nanoarchaeum equitans*' and 'Nanoarchaeota' strain Nst1; Podar *et al.*, 2013), it appears to possess pathways allowing for the independent production of ATP and the biosynthesis of multiple key metabolites. To examine whether the observed metabolic capacities within Cand. IA are because of a reductive

evolutionary process of gene loss from a metabolically versatile ancestor, or to gene acquisition by a metabolically limited ancestor, we analyzed the occurrence and prevalence of HGT events within Cand. IA genome. Because of the limited genomes available representing the DPANN superphylum, identification of incidents of HGT from an archaeal donor was not feasible. Therefore, our analysis was limited to the identification of genes having non-archaeal origins within Cand. IA genome.

Cand. IA genome showed a higher percentage of proteins (25.4%,  $n=305$ ) with apparent bacterial Blastp first hits compared with all other archaeal genomes examined (those ranged from 1.44% in '*Nanoarchaeum equitans*' to 16.9% in *Candidatus Micrarchaeum acidiphilum* ARMAN2; Figure 2a). A disproportionately large percentage (116 out of 196) of metabolism-related genes in Cand. IA genome were of apparent bacterial origin, with the majority of these metabolic genes (86 out of 116) involved in biosynthetic pathways (Figure 2b). Maximum likelihood analysis of the phylogenetic affiliation of 21 'Diapherotrites' anabolic proteins confirmed their putative bacterial origin (Supplementary Figure S1). Detailed analysis of the impact of HGT on every metabolic pathway examined is summarized in Table 2 and Supplementary Table S3. Out of 26 metabolic pathways examined, 21 showed evidence of HGT with at least one gene in the pathway with bacterial Blastp first hit. In some cases, an entire metabolic pathway was completely bacterial in origin, for example, biosynthesis of threonine, histidine, arginine and proline. Within other pathways, a fraction of the genes were bacterial in origin; but these genes appear to mediate the critical/defining steps of the pathway, for example, PHB depolymerase in PHB degradation pathway and phosphoenolpyruvate synthase in gluconeogenesis. It is notable that many of the metabolic pathways affected by HGT are absent from '*Nanoarchaeum equitans*', the model Nanoarchaeal obligate endosymbiont (Table 2 and Supplementary Table S3). Phylogenetic analysis suggests multiple bacterial phyla as potential gene donors to the Cand. IA genome (Table 2), although within HGT acquired genes, overlapping genes as well as genes with multiple copies usually appear to have the same bacterial donor.

Collectively, the acquired genes enabled Cand. IA to: (1) synthesize alanine, threonine, arginine, proline, histidine and aromatic amino acids, purine and pyrimidine nucleotides and the cofactor pyridoxal phosphate, (2) to convert pterin to folate, (3) to uptake and activate thiamine, (4) to break down proteins extracellularly and potentially uptake the resulting oligopeptides and amino acids and (5) to utilize alanine and valine as possible C and energy sources, and to depolymerize PHB. These results demonstrate that cross-kingdom HGT events represent an important mechanism that contributes to enhancing the metabolic capacities of Cand. IA.

**Table 2** Metabolic features and potential origins of metabolic genes in *Cand. IA* genome

Pathway	No. of HGT/total <sup>a</sup>	Potential bacterial phyla donor(s)	Nanoarchaea <sup>b</sup>
<i>I. Biosynthetic pathways</i>			
<i>A. Pathways entirely bacterial in origin</i>			
<i>1. Amino acids</i>			
a. Gly from Ser	1/1	Deferribacter	0/1
b. Ala from pyruvate	1/1	Thermodesulfobacteria	0/1
c. Glu from Ala and $\alpha$ -KG	1/1	Thermodesulfobacteria	0/1
d. Thr from Asp (missing 1)	4/4	Bacteroidetes, Acidobacteria, Spirochaetes	0/4
e. His from PRPP	8/8	Acidobacteria, Synergistetes, Proteobacteria, Firmicutes, Aminicenantes, and Parcubacteria	2/9
f. Arg from carbamoyl-P and Asp	6/6	Thermotoga, Firmicutes, Spirochaetes, Bacteroidetes and Cyanobacteria	1/6
g. Pro from Glu	3/3	Firmicutes and Aquificae	0/3
<i>2. Cofactors</i>			
a. B6	2/2	Armatimonadetes, and CD JS-1	0/2
b. Folate and 1C pool	7/7	Firmicutes, Thermotoga, Proteobacteria, and Deferribacter	0/7
<i>B. Pathways partially bacterial in origin</i>			
<i>1. Amino acids</i>			
a. Asn from Asp	2/5	Firmicutes and Aquificae	3/5
b. Aromatic amino acids from 7-phospho-2-dehydro-3-deoxy-D-arabino-heptonate			
i. Trp	8/12	Proteobacteria, Thermodesulfobacteria, Planctomycetes,	2/12
ii. Phe	7/9	Nitrospira, Firmicutes and Hydrogenedentes	3/9
iii. Tyr			
<i>2. Cofactors</i>			
a. Thiamine	3/4	Firmicutes and Proteobacteria	1/4
b. Riboflavin	2/6	Firmicutes	0/6
c. Nicotinate and nicotinamide	2/6	Firmicutes and Proteobacteria	0/6
d. Pantothenate and CoA	1/4	Aminicenantes (CD-OP8)	0/4
<i>3. Nucleotides</i>			
a. Purine from ribose-1-P	10/12	Planctomycetes, Proteobacteria, Firmicutes, Lentisphaera, and Omnitrophica	1/12
b. Pyrimidine from glutamine and PRPP	11/18	Bacteroidetes, Proteobacteria, Cyanobacteria, Firmicutes, Planctomycetes, and Omnitrophica	2/18
<i>4. Carbohydrates</i>			
Gluconeogenesis to fructose-6-P	6/13	Firmicutes, Bacteroidetes, Proteobacteria, and Parcubacteria	9/13
<i>C. Pathways entirely archaeal in origin</i>			
a. Ser from 3-P glycerate	0/3	NA	2/3
b. Gln from Glu	0/6	NA	1/6
c. Asp from oxaloacetate	0/1	NA	0/1
<i>II. Catabolic pathways (partially bacterial in origin)</i>			
1. PHB degradation	2/6	Actinobacteria and Proteobacteria	2/6
2. Ribose degradation	5/9	Thermodesulfobacteria, Acidobacteria, Firmicutes, Bacteroidetes, and Parcubacteria	4/9
3. Peptidases	3/5	Actinobacteria and Firmicutes	1/5
4. Potential ETS	2/12	Proteobacteria and Firmicutes	5/12

Abbreviations: *Cand. IA*, *Candidatus 'Iainarchaeum andersonii'*; HGT, horizontal gene transfer; PHB, polyhydroxybutyrate; PRPP, 5-phospho- $\alpha$ -D-ribose 1-pyrophosphate.

<sup>a</sup>Number of HGT candidates/total number of genes in the pathway.

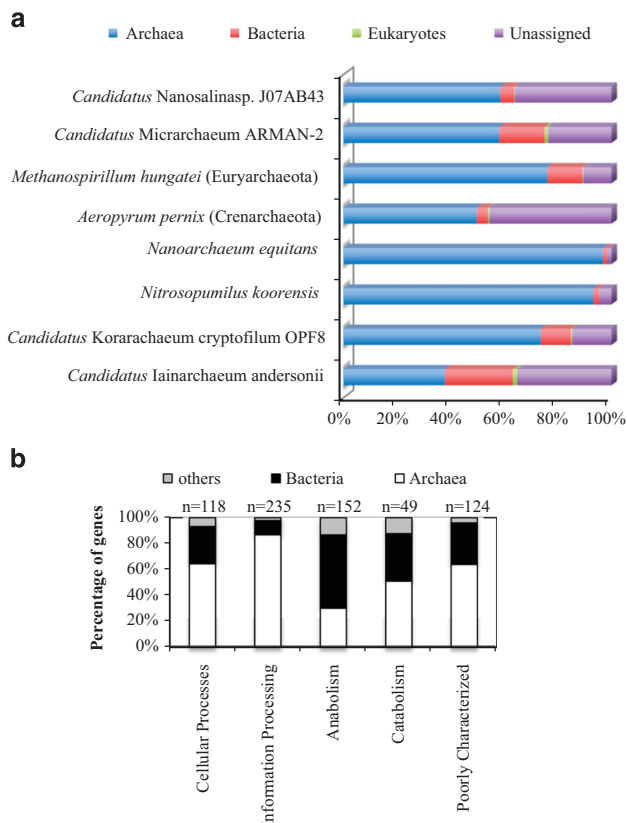
<sup>b</sup>Number of genes/total number of genes in the pathway with homologs in the published Nanoarchaeal genomes.

### Genome architecture and COG distribution patterns in *Cand. IA* genome

Various genomic features and COG distribution patterns in microbial genomes have successfully been correlated to putative trophic lifestyles, for

example, oligotrophy, copiotrophy (Giovannoni *et al.*, 2005; Lauro *et al.*, 2009; Swan *et al.*, 2013) as well as obligate symbiosis (Moran and Wernegreen, 2000; Shigenobu *et al.*, 2000; Akman *et al.*, 2002; Tamas *et al.*, 2002; Moran *et al.*, 2003;





**Figure 2** (a) Phylogenetic distribution of non-self Blastp first hits of *Candidatus Iainarchaeum andersonii* proteins compared with other archaeal phyla representatives. (b) Phylogenetic distribution at the domain level of non-self Blastp first hits of *Candidatus Iainarchaeum andersonii* proteins classified by metabolic category in the X axis. Total number of proteins belonging to each category are shown above each column.

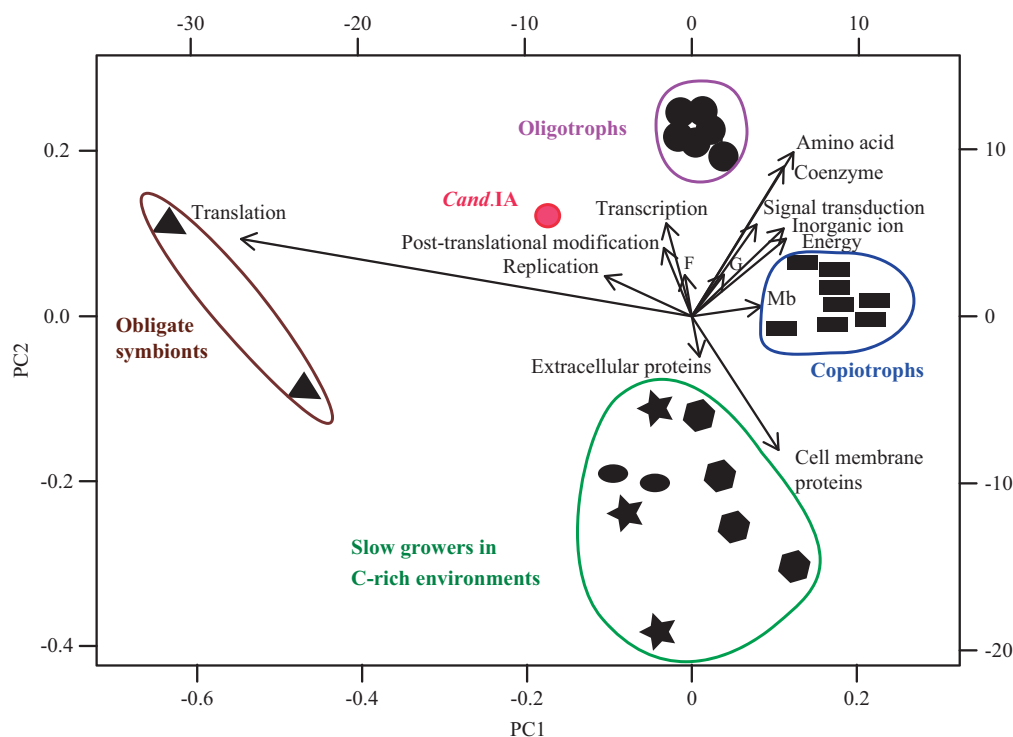
Waters *et al.*, 2003; Moya *et al.*, 2008; Nikoh *et al.*, 2011; Hendry *et al.*, 2013; Podar *et al.*, 2013). In an effort to decipher the putative trophic lifestyle of Cand. IA, we used PCA to compare Cand. IA genome with those of 19 other archaeal genomes (Supplementary Table S2) encompassing obligate oligotrophs, obligate archaeal symbionts, fast-growing copiotrophs as well as the slow-growing archaeal copiotrophs (marine groups MCG and Thermoplasmatales MBG-D group thriving in C-rich ocean sediments). PCA biplot (Figure 3) showed that, in general, genomes clustered according to their trophic lifestyle into four major groups: fast-growing copiotrophs, slow-growing copiotrophs, obligate oligotrophs and obligate symbionts, with the later being highly divergent and clustering away from other genomes. This was expected as archaeal symbionts exhibit a dramatic reduction in metabolism-related and an expansion in information processing-related gene families (Waters *et al.*, 2003; Podar *et al.*, 2013).

Cand. IA genome did not cluster with any of the above groups in the PCA biplot, but rather showed a distinct position between the obligate nanoarchaeal symbionts and the archaeal oligotrophs.

This position is a reflection of the overrepresentation of replication-related, post-translational modification-related and nucleotide metabolism-related proteins compared with other genomes, as well as the overrepresentation of translation-related proteins compared with all other genomes with the exception of ‘Nanoarchaeota’. The position of Cand. IA in the same quadrant with archaeal symbionts and oligotrophs is a reflection of the shared streamlining genomic characteristics, for example, smaller genome size and significantly lower percentage of cell membrane proteins, when compared with copiotrophs. However, salient differences exist between Cand. IA and typical oligotrophs and obligate symbionts. Compared with ‘*Nitrosopumilus maritimus*’ genome (a model archaeal oligotroph), Cand. IA genome has an overrepresentation of replication, translation, transcription, nucleotide metabolism, intracellular trafficking and secretion and cell wall biogenesis COGs as well as proteins destined to the cell wall, and an underrepresentation of signal transduction, defense mechanisms, energy, amino acid, coenzyme, inorganic ion and secondary metabolism COGs as well as transporters and extracellular proteins (Figure 3 and Supplementary Table S2). Similarly, compared with ‘*Nanoarchaeum equitans*’ genome (a model archaeal obligate symbiont), Cand. IA genome has a larger genome, an overrepresentation of energy, amino acid, nucleotide, carbohydrate, coenzyme, lipid and inorganic ion metabolism, cell wall biogenesis and signal transduction COGs, and an underrepresentation of translation, replication and post-translational modification COGs. Interestingly, many of these defining features between archaeal symbionts and Cand. IA could be mediated by the acquisition of genes via HGT as described above. The disparate position of all DPANN genomes analyzed is striking, and underscores the high level of diversity in genomic architecture, metabolic potential and trophic lifestyle within the DPANN superphylum (Rinke *et al.*, 2013).

#### Global distribution of the ‘*Diapherotrites*’

The 16S rRNA genes obtained from the three available ‘*Diapherotrites*’ SAG genome assemblies were shorter (average length is 1312bp) than other archaeal counterparts, mainly because of the absence of bases corresponding to 1–20 and 1381–1540 in *Methanobacterium formicicum* (Gutell *et al.*, 1985). Therefore, universal 16S rRNA gene primers targeting these regions (for example, A1F, U1406R and U1510R) (Baker *et al.*, 2003) would theoretically fail to identify ‘*Diapherotrites*’ members. Furthermore, within these shortened 16S rRNA genes, mismatches to almost all known archaeal-specific or universal 16S rRNA gene primers were identified in all three available ‘*Diapherotrites*’ SAG genome assemblies, with the exception of 109F, 515F and 534R (Table 3).



**Figure 3** PCA biplot of the genomic features and COG category distribution in the genomes compared. Genomes are represented by symbols according to their trophic lifestyle. A list of genomes, as well as features used in this analysis, and trophic lifestyles are presented in Supplementary Table S2. Arrows represent genomic features or COG categories used for comparison. The arrow directions follow the maximal abundance, and their lengths are proportional to the maximal rate of change between genomes. The first two components explained 75% of variation. Obligate oligotrophs of the Thaumarchaeota (depicted by circles) clustered together because of abundance of amino acid, nucleotide and coenzyme metabolism-related as well as post-translational modification and transcription-related proteins, the copiotrophs (depicted by rectangles) clustered together mainly because of their large genome sizes and a higher percentage of membrane proteins relative to other groups, and the slow-growing copiotrophs of the Thermoplasmatales (depicted by hexagons) clustered together because of the expansion of membrane as well as extracellular proteins consistent with previous reports of a higher percentage of membrane transporters and extracellular peptidases in those genomes (Lloyd *et al.*, 2013). Parvarchaeota genomes (depicted by stars) and Nanoarchaeota genomes (depicted by ovals) also clustered close to the slow-growing copiotrophs. Finally, the two obligate symbionts of Nanoarchaeota (depicted by triangles) clustered together away from all other genomes mainly because of expansion of translation-related proteins. *Candidatus Iainarchaeum andersonii* genome is represented by a red circle and has an intermediate position in the plot. F, COG category nucleotide metabolism; G, COG category carbohydrate metabolism; Mb, genome size.

With mismatches to known primers, ‘Diapherotrites’ sequences would theoretically be missed in cultivation-independent PCR-based surveys. Indeed, comparison of ‘Diapherotrites’ to Sanger-generated, near-full-length 16S rRNA genes deposited in GenBank nr database failed to identify any ‘Diapherotrites’-related 16S rRNA gene sequences. In addition, within a 16S rRNA data set generated using primer pair 926wF and 1392R from the same sample from which the three ‘Diapherotrites’ SAGs were obtained, no ‘Diapherotrites’ 16S rRNA gene sequences were amplified (Supplementary Table S4). Furthermore, within a collection of 31 972 882 archaeal next-generation sequences spanning 58 habitats and 775 data sets, only 66 sequences were confidently assigned to the ‘Diapherotrites’ phylum from 3 different studies, where they were identified in a paddy soil (Feng *et al.*, 2013), three distinct soils (Portillo *et al.*, 2013) and wastewater treatment plant (Vishnivetskaya *et al.*, 2013).

Finally, we used phylogenetic anchoring to identify the presence of members of the ‘Diapherotrites’

in publicly available metagenomic data sets ( $n = 893$ ). The ‘Diapherotrites’ were only identified in a handful of environments (11 out of 893 analyzed). These include Amazon forest soil, mangrove sediment on Isabella Island, Sakinaw lake, Etoliko lagoon sediment and Kolumbo Volcano red mat (Supplementary Figure S2). Within these studies, the ‘Diapherotrites’ were always identified as an extremely minor fraction of the community ( $< 0.006\%$  of anchored metagenomic reads).

## Discussion

In this study, we present a detailed analysis of the metabolic capabilities and genomic features of three SAGs belonging to the recently proposed archaeal phylum ‘Diapherotrites’, as well as a survey of the putative distribution of members of this phylum using database-mining approaches. Our detailed genomic analysis of ‘Diapherotrites’ SAG *Cand. IA* uncovers evidence for genome streamlining;

**Table 3** Mismatches of 'Diapherotrites' 16S rRNA gene to universal archaeal primers<sup>a</sup>

Primer name	Primer sequence (5'–3')	Sequence in <i>Diapherotrites</i> (5'–3')
A109F	ACKGCTCAGTAACACGT	ACKGCTCAGTAACACGT
A333F	TCCAGGCCCTACGGG	<b>CC</b> TAGCCCTAMGGG
AB341F	CCTACGGGRSGCAGCAG	CCTAMGGGRTGCAGCAG
A344F	ACGGGGTGCAGCAGGCGGA	AMGGGGTGCAGCAG <b>KYR</b> SGA
U515F	GTGCCAGCMGCCGCGGTAA	GTGCCAGCMGCCGCGGTAA
U519F	CAGCMGCCGCGGTAATWC	CAGCMGCCGCGGTAATWC
UA571F	GCYTAAAGSRICCGTAGC	GCYTAAAGSR <b>IYY</b> GTAGC
UA751F	CCGACGGTGAGRGRYGAA	CTGACGGTGAG <b>RRR</b> YGAA
AB779F	GCRAASSGGATTAGATACCC	GCRA <b>CS</b> SGGATTAGATACCC
EB787F	ATTAGATACCCCTGGTA	ATTAGATACCCGGT <b>T</b> A
AB787F	ATTAGATACCCGGGTA	ATTAGATACCCGGT <b>T</b> A
Ab789F	TAGATACCCSSGTAGTCC	TAGATACCC <b>S</b> TTAGTCC
AB906F	GAAACTTAAAKGAATTG	GAAACTTAA <b>W</b> KGAATTG
A1040F	GAGAGGWGGTGCATGGCC	GAGAGGWGYTGTATGGYC
U1053F	GCATGGCYGCGTCAG	GTATGGCY <b>A</b> YK <b>K</b> CAG
A1098F	GGCAACGAGCGMGACCC	GGCAACGGCGCGMGACCY
AB127R	CCACGTGTTACTSAGC	CBACGTGTTACTSAGC
A348R	CCCCGTAGGGCCYGG	CCCC <b>K</b> TAGGGCTAGG
U534R	GWATTACCGCGCKGCTG	GTATTACCGCGCGCKGCTG
U529R	ACCGCGGCKGCTGGC	ACCGCGCGCKGCTGGC
AB909R	TTTCAGYCTTGCGRCCGTAC	TTT <b>C</b> AC <b>Y</b> CTTGCGR <b>G</b> YRTAC
A927R	CCGCCAATTCCTTAAAGTTTC	CCGCCAATTC <b>C</b> WTTAAAGTTTC
A926R	CCGTCAATTCCTTTRAGTTT	CCGCCAATTC <b>C</b> WTTTRAGTTT
A934R	GTGCTCCCGGCCAATTCCT	GTG <b>M</b> G <b>Y</b> CCCGGCCAATTC <b>C</b> W
A976R	YCCGGCGTTGAMTCCAATT	YCYGGCGTT <b>G</b> TRTCC <b>R</b> ATT
A1115R	GGGTCTCGCTCGTTG	RGGTCTCGCYCGTTG
UA1204R	TTMGGGGCATRCIKACCT	TTMGGG <b>C</b> YATRCIKAYCT

Mismatches in 'Diapherotrites' sequences are shown in bold.

<sup>a</sup>Using the 16S rRNA gene sequences of the three *Diapherotrites* single amplified genomes (SAGs).

prevalence of HGT events, especially in metabolism-related genes; limited catabolic capabilities with only few substrates that could potentially be utilized for ATP production; and a limited representation of members of this phylum in amplicon-generated and metagenomic data sets.

Many of the genomic streamlining features observed in *Cand. IA* genome, such as small genome size, small intergenic regions, low incidences of gene duplication and low number of rRNA operons, have been associated with specific trophic lifestyles, mainly oligotrophy and obligate symbiosis (Giovannoni *et al.*, 2005; Lauro *et al.*, 2009; Walker *et al.*, 2010; Grote *et al.*, 2012; Swan *et al.*, 2013), where they appear to be a reflection of the accessibility of nutrients, as well as the occurrence of genetic drift in obligate symbionts (Mira *et al.*, 2001; Wernegreen, 2002; Giovannoni *et al.*, 2005; Oakeson *et al.*, 2014). However, detailed comparative analysis of the metabolism and genomic features of *Cand. IA* revealed salient differences when compared with the genomes of model archaeal obligate symbionts and oligotrophs.

Compared with the genome of the model archaeal obligate symbiont '*Nanoarchaeum equitans*', *Cand. IA* genome possesses multiple catabolic abilities that allow for the production of ATP from few substrates (valine, alanine, aspartate, ribose and PHB) (Figure 1, Table 2, and Supplementary Text). Such capabilities are completely absent from '*N. equitans*' genome (Waters *et al.*, 2003).

More importantly, *Cand. IA* possesses an anabolic machinery that allows for the biosynthesis of multiple amino acids, nucleotides and cofactors (Figure 1, Table 2 and Supplementary Text); a feature that is absent in '*N. equitans*' because of its dependence on its host for supplying such metabolites (Waters *et al.*, 2003).

In contrast to archaeal oligotrophs that have numerous transport capabilities, a well-developed essential biosynthetic machinery as well as complete central metabolic pathways (Walker *et al.*, 2010; Nunoura *et al.*, 2011), *Cand. IA* exhibits lower transport capabilities (Supplementary Table S2), higher level of auxotrophy and incomplete and/or less developed pathways (for example, respiratory chain, tricarboxylic acid cycle and pentose phosphate pathway). Furthermore, the catabolic capabilities of *Cand. IA* appear to be geared toward utilizing substrates that are more common in non-oligotrophic habitats. For example, ribose, being a lysis product of RNA, is presumably more available in non-oligotrophic environments characterized by higher rates of cell turnover and lysis. Indeed, both the transporters and the carbohydrate utilization patterns of the SAR11 clade comprising the oligotrophic ocean bacteria suggest the inability to take-up and utilize carbohydrates (including ribose) as a carbon and energy source (Jiao and Zheng, 2011). Similarly, PHBs are storage molecules produced by several bacterial species in response to either an excess of carbon or a limitation of another nutrient,

for example, nitrogen or phosphorous (Jendrossek and Handrick, 2002), and hence would be expected to exist in C-rich environments (Jendrossek and Handrick, 2002).

Analysis of the global ecological distribution of members of the ‘Diapherotrites’ identified its presence in only a few environments (Supplementary Figure S2). However, this observed pattern of paucity of ‘Diapherotrites’ sequences in either amplicon-generated or metagenomic data sets could be influenced by the mismatches identified to the most commonly utilized 16S rRNA gene primers (Table 3), or the limited number of ‘Diapherotrites’ reference sequences available to serve as substrates for phylogenetic anchoring analysis, respectively. Nevertheless, examination of the origin and trophic status of habitats where the ‘Diapherotrites’ were identified suggest their preference to non-oligotrophic environments (for example, microbial mats, high productivity forests, wastewater treatment plants (Vishnivetskaya *et al.*, 2013) and soils (Feng *et al.*, 2013; Portillo *et al.*, 2013)).

We argue that the observed metabolic capacities, genomic features and ecological distribution as well as the observed high proportion of genes involved in cross-kingdom HGT (Table 2 and Supplementary Table S3) could be explained by a conceptual model where gene acquisition plays an important role in shaping the evolutionary history of Cand. IA. Specifically, we argue that this acquisition process is mediating the putative transition of Cand. IA from a symbiotic ancestor with a streamlined genome and extremely limited metabolic capabilities to a free-living microorganism, capable of ATP production (although from a limited number of substrates), as well as biosynthesis of multiple amino acids, nucleotides and cofactors, although it remains auxotrophic to other several cellular building blocks. Indeed, most of the key differences in genome architecture between ‘*N. equitans*’ and Cand. IA (Figure 3) could be brought about by the presence of additional genes of apparent bacterial origin in the genome assembly. Theoretically, removal of genes of bacterial origin from the Cand. IA genome would produce a genome assembly with features and metabolic capacities very similar to the genome ‘*N. equitans*’ (Supplementary Figure S3).

Although the role of HGT in conferring specific capabilities to recipient prokaryotic species, for example, antibiotic resistance or heavy metal resistance (Andam *et al.*, 2011; Navarro *et al.*, 2013), has long been recognized, the impact of HGT on prokaryotic evolutionary history and its potential role in organismal transition to new habitats and lifestyles has received less attention. The acquisition of bacterial genes was recently proposed as a driver of Halobacteriales evolution from a methanogenic ancestor (Nelson-Sathi *et al.*, 2012). Within the eukaryotes, HGT has been shown to be important in the development of thermoacidophily and subsequent adaptation of the red algae *Galdieria*

*sulphuraria* to hot acidic habitats (Qiu *et al.*, 2013; Schönknecht *et al.*, 2013), as well as the adaptation of gut fungi (Neocallimastigomycota) to the strict anaerobic, eutrophic and plant biomass-rich habitat in the herbivorous gut (Youssef *et al.*, 2013). Additional research to provide a more detailed understanding of the impact of such processes on microbial (especially prokaryotic) evolution is certainly warranted.

## Conflict of Interest

The authors declare no conflict of interest.

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