Genome Sequence Analysis and Identification of Genes Associated to Pesticide Degradation from Enterobacter cloacae Strain MR2

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ABSTRACT

Today's burning problem in the world is pesticide residues in foods. To overcome this problem, nineteen chlorpyrifos-degrading bacteria were isolated from soil of adjoining area of pesticide manufacturing industries located in Gujarat, India. The strain CPD-12 (MR2) degraded highest chlorpyrifos among the other strains isolated from different sites, i.e. Up to 500 ppm in 30 hrs. And hence was selected further for whole genome sequencing. This strain showed maximum similarity to members of the order Enterobacteriales and was closest to Enterobacter cloacae of this group. The genome sequence of strain Enterobacter cloacae MR2 consisted of a circular 4,758,062bp chromosome with a 55.1% G+C value, 5571 protein coding genes, 16rRNA and 72 tRNAs. The genome annotation and functional characterization of the strain MR2 provided insights into various genetic processes involved in the degradation of several pesticides and detoxification of toxic compounds. The genome of MR2 was also compared with Enterobacter cloacae subsp cloacae ATCC 13047 and Enterobacter sp. 638 which showed the presence of genes for the pesticide degradation as in ATCC 13047 and also had genes to promote plant growth as in Enterobacter sp. 638.

K E Y W O R D S
Whole genome sequencing, chlorpyrifos degradation, Enterobacter cloacae MR2, Draft genome

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Introduction

Organophosphates (OPs) pesticides are highly toxic chemical pesticide that exhibit broad-spectrum activity against insects and accounts for about 38% of the total pesticides used globally for agricultural crops. Continuous and excessive use of OPs has caused not only nerve (this class of pesticide has acute neurotoxicity due to their ability to suppress acetyl-choline-esterase) and muscular diseases in human and animals but also have contaminated ecosystems in different parts of the world (Zhang et al., 2008).

Chlorpyrifos (O, O-diethyl O-3, 5, 6-trichloropyridin-2-yl phosphorothioate) as an active ingredients a broad spectrum organophosphorus insecticide, most widely used for pest control (Cho et al., 2002). It has been widely used for aerial application to control surface feeding insects (Dhawan and Simwat, 1996; Gupta et al., 2001; Sasikala et al., 2012) and also applied to soil for root
damaging insect larvae (Bhatnagar and Gupta, 1992; Rouchaud et al., 1991; Davis et al., 1976). Pesticides and their degradation products generally get accumulated in the soil and influence not only the population of various groups of microbes, but also their biochemical activities like nitrification, ammonification, decomposition of organic matter and nitrogen fixation (Agnihotri et al., 1981; Faldu et al., 2014). In soil, chlorpyrifos may remain biologically active for periods ranging from days to months. Dosage rates, soil type, soil moisture and organic matter, content, temperature and insecticide formulation are among the factors which influence the biological persistence (Read 1976; Tashiro et al., 1978) it is moderately persistent in nature as its residues were detected in soil even after 3 months of application and hence causes potential environmental hazards (Chapman et al., 1984).

Microorganisms play an important role in degrading synthetic chemicals in soil (Alexander, 1981). They have the broad capacity to utilize almost all natural and some synthetic compounds as their sole carbon and energy source. Chlorpyrifos degrading bacteria can be used either directly or indirectly, for the bioremediation of chlorpyrifos contaminated soils. Till now, various genes, such as opd (organophosphate-degrading) and mpd (methyl parathion degrading) and several enzyme systems have been identified which were found to be involved in degradation of certain organophosphates (Serdar, 1982; Mulbry et al., 1986; Horne et al., 2002; Yang et al., 2006; Cui et al., 2001; Parakhia et al., 2014).

In the present study, chlorpyrifos degrading bacteria were isolated from various pesticide contaminated sites and were screened for their chlorpyrifos degradation capability through High Performance Liquid Chromatography (HPLC). The most efficient bacterium Enterobacter cloacae stain MR2 was sequenced for complete genome. The genome of E. cloacae MR2 was characterized for identification of genes responsible for the degradation of chlorpyrifos and was also compared with Enterobacter cloacae subsp cloacae ATCC 13047 and Enterobacter sp. 638 for synteny.

Materials and Methods

Isolation and screening of chlorpyrifos degraders

Soil samples were collected from five different sites which were contaminated regularly with the pesticides from Gujarat, India (Table 1). Out of 45 strains initially isolated, 19 were screened out with the ability to degrade 50-500 ppm chlorpyrifos by Shake flask method and were quantified by HPLC among 19 strains, CPD-12 was found to be most efficient degrader with the ability to degrade 500 ppm within 30 hrs was selected for the genome sequence analysis.

Genome sequencing

For genome sequencing, DNA of Enterobacter cloacae stain MR2 was isolated using Phenol-Chloroform method (Sambrook et al., 1989). The DNA concentration and purity was determined using Picodrop PET01 (Picodrop Ltd., Cambridge, U.K).

The DNA was enzymatically fragmented to construct a library of 260 bp, which was further used for template preparation. Sequencing was carried out using Ion Torrent Personal Genome Machine (PGM™) from Life Technologies, at Department of Biotechnology, Junagadh Agricultural University, Junagadh, India as per the manufacture's guidelines.
Gene prediction and annotation

Raw reads of the sequence were processed for the quality control through default plug-in in Ion Torrent Software Server (FastQC). The quality reads were assembled in MIRA v 3.4.1 by using Smith-Waterman algorithm (Chevreux et al., 2000). Contings were ordered through the tool Mummer (Kurtz et al., 2004) and were aligned with reference genome E. cloacae ATCC 13047 and Enterobacter sp. 638 using Mauva (Darling et al., 2010) software. Putative coding sequences (CDS) were initially identified by RAST automated annotation software (Aziz et al., 2008; Overbeek et al., 2014) followed by Magnifying Genome annotation platform (MaGe) (http://www.genoscope.cns.fr/agc/mage/). All CDS identified were manually reviewed, and false CDS were flagged as “artifact”. The remaining CDS were then submitted to automatic functional annotation via BLAST searches against the UniProt databank in order to determine significant homology. Circular chromosomal map of E. cloacae MR2 with annotated genes/CDS was constructed using CGView (Stein et al., 2001). Core and Pan Genome analysis of E. cloacae MR2 with E. cloacae ATCC 13047 and Enterobacter sp. 638 was analyzed by MaGe-Microscope Pan Genome Analysis interface (Vallenet et al., 2006).

Results and Discussion

Characterization of bacterial strain

Soil samples collected from five chlorpyrifos contaminated sites of Gujarat, India resulted in the isolation of 45 stains. Out of 45, 19 strains were able to degrade chlorpyrifos (Table 2) and among them, CPD-12 was found to be most efficient degrade with the ability to degrade 500 ppm within 30 hrs.

Genome de Novo assembly

Whole genome sequencing of Enterobacter cloacae MR2 was carried out using Ion Torrent (PGM) whole genome sequencer (Life Technologies) at the Department of Biotechnology, JAU, Junagadh. A total of 549,959 reads with an average length of 176 bp and have coverage of the 18.06X. Initial quality check of raw data was performed through FASTQC and reads were filtered based on base quality and length (Fig. 1). Quality reads were assembled by MIRA which resulted in 230contigs with longest contig of 177,145bp and N95 of 7,065 bp (Table 3-4).

Assembled genome was submitted to an automated annotation tool RAST (Rapid Annotation using Subsystem Technology), which provides high quality genome annotations for bacterial and archaea genomes. RAST indicated E. cloacae subsp cloacae ATCC 13047 (score 500) and Enterobacter sp. 638 (score 452) as the closest members of E. cloacae MR2. The 'neighbor' score in RAST was estimated via quick and dirty” ad hoc heuristic method which is based on the number of times that the 'neighbor' genome was the top hit in BLAST against the candidate (in this case E. cloacae MR2) from the set of "unique" genes within the query genome. A higher score suggested that the two genomes are likely to be metabolically similar. The comparative profile of the MR2 RAST distribution (Fig. 2a) covers 4068 subsystems compare to 3909 of ATCC 13047 and 3564 of strain 638. MR2 codes highest for the metabolism of carbohydrates and amino acids followed by membrane transport mechanism and production of cofactor, vitamins, prosthetic groups, pigments. Complete genome of Enterobacter cloacae subsp cloacae ATCC 13047 (ref: NC_014121.1) and Enterobacter sp. 638 genome from NCBI (ref:
NC_009436.1) were used for comparative and synteny analysis (Fig. 3). In the synteny map the pattern of the arrangement of the blocks indicated variation with reference genome. The space between two blocks indicates the gap region, which was not found similar in the referred genome, may be probable horizontally transferred regions.

**Genome annotation**

Genome annotation is the process of attaching biological information to sequences. It includes predicting genes function, structure, coding regions and ORFs. Genome annotation of *E. cloacae* MR2 predicted various genes involved in various stress response(s) as well as genes involved in resistance to antibiotics and toxic compounds as indicated by RAST analysis (Fig. 5).

Total 64 genes that are involved in multi-drug resistance, multiple antibiotic resistance, resistance to Fluoro-quinolones, copper homeostasis and tolerate to heavy metals like Copper, Cobalt, Arsenic, Zinc, Cadmium etc. Were identified. Above this, annotation also revealed 32 stress responsive genes that coded for universal stress response protein family, phage shock proteins and genes involved in various processes like carbon starvation, sugar phosphate stress regulation etc. Along with these, genes that take part in other stress responses were also identified which included 59 genes for oxidative stress, 9 for desiccation stress and 24 for osmotic stress / Osmo-regulation. Thirty one gene that is responsible for detoxification like Nudix proteins (nucleoside triphosphate hydrolases) which are activated in plant defense response, a family of versatile, widely distributed housekeeping enzymes, housekeeping nucleoside triphosphate pyrophosphatases, genes involved in tellurite resistance and chromosomal determinants etc. Were identified. The annotation also indicated the presence of genes for phosphorus metabolism, sulfur metabolism, metabolism of aromatic compounds, nitrogen metabolism, protein metabolism, potassium metabolism and iron acquisition and metabolism in the genome of strain MR2.

Enzymes responsible for catabolism of organophosphate compound such as Inorganic Pyrophosphatase (EC 3.6.1.1), Phosphonoacetaldehyde hydrolase (EC 3.11.1.1), 3-ketoacyl-CoA thiolase (EC 2.3.1.16), Salicylate hydroxylase (EC 1.14.13.1), Catechol 1,2-dioxygenase (EC 1.13.11.1), 1H-3-hydroxy-4-oxoquinodimine 2,4-dioxygenase, Catechol 2,3-dioxygenase (EC 1.13.11.2), Gentisate 1,2-dioxygenase (EC 1.13.11.4) and Monoamine oxidase (EC 1.4.3.4) were also identified during the process of annotation. Metabolic Reconstruction of *Enterobacter cloacae* MR2 and *Enterobacter* sp. 638 allowed the comparison of functioning parts of two organisms (Table.5; Fig.2b).

It provided a list of all genes which were associated with a subsystem in the respective organism. Genes for stress responsive (142), phosphorus metabolism (41) and sulphur metabolism (54) were found to be common in both genomes. While comparing with *Enterobacter cloacae* subsp *cloacae* ATCC 13047, MR2 cluster of orthologous genes (COG) categories indicated highest distribution for the general function prediction (718), Amino acid transport and metabolism (657), Carbohydrate transport and metabolism (596) while ATCC 13047 revealed 626,500 and 463 CDS respectively (Table 3b).

The BLAST map of *Enterobacter cloacae* MR2 with the other genomes present in the microscope platform software indicated *E. cloacae* ATCC 13047 as the nearest genome (Fig. 4), followed by *E. hormaechei* ATCC 49162, *E. concorogenus* ATCC 35316, *Enterobacter* sp. 638 and *E. aerogenes* KTCC 2190.
### Table 1 Locations of screening of chlorpyrifos degrading bacteria

| Sr. No. | Name of CPD isolate | Industry | GPS Coordinates | Area       | District   |
|---------|---------------------|----------|-----------------|------------|------------|
| 1-4     | CPD-1, CPD-2, CPD-3, CPD-4 | Near Pioneer Agro Industry | Latitude: 23.070887 | Longitude: 72.671289 | Ahmedabad G.I.D.C. | Ahmedabad |
| 5-13    | CPD-5, CPD-6, CPD-7, CPD-8, CPD-9, CPD-10, CPD-12, CPD-13 | United Phosphorus Limited | Latitude: 21.618039 | Longitude: 73.022817 | Ankleshwar G.I.D.C. | Bharuch  |
| 14-17   | CPD-14, CPD-15, CPD-16, CPD-17 | Near GIDC, Kadi, Gujarat, India | Latitude: 23.29042 | Longitude: 72.36219 | Kalol G.I.D.C. | Ahmedabad |
| 18      | CPD-18 | Field Collection Ivnagar | Latitude: 21.477184 | Longitude: 70.43203 | Field Collection Ivnagar & Vadla | Junagadh |
| 19      | CPD-19 | Field Collection Vadla | Latitude: 21.477991 | Longitude: 70.40041 |

### Table 2 Concentration of standard chlorpyrifos at different incubation period

| Sr. No. | Name of CPD isolates | Concentration of standard chlorpyrifos at different incubation period (mg/l) |
|---------|---------------------|--------------------------------------------------------------------------|
|         |                     | 6 hr | 12 hr | 18 hr | 24 hr |  |
| 1       | CPD-1               | 248  | 195.47 | 139.86 | 90    |
| 2       | CPD-2               | 264.18 | 223.99 | 188.99 | 151.43 |
| 3       | CPD-3               | 238.5 | 189.04 | 149.46 | 124.15 |
| 4       | CPD-4               | 241.72 | 186.07 | 102    | 66.56  |
| 5       | CPD-5               | 261.5 | 217.94 | 187.09 | 145.24 |
| 6       | CPD-6               | 254.29 | 214.33 | 159.55 | 123.68 |
| 7       | CPD-7               | 252.75 | 203.6  | 154.04 | 110.79 |
| 8       | CPD-8               | 262.1 | 229.2  | 173.25 | 131.1  |
| 9       | CPD-9               | 246.13 | 193.02 | 162.92 | 102    |
| 10      | CPD-10              | 245.24 | 191.57 | 110.25 | 75.64  |
| 11      | CPD-11              | 243.15 | 179.05 | 130.94 | 87.71  |
| 12      | CPD-12              | 234.17 | **163.66** | **80.26** | **22**  |
| 13      | CPD-13              | 254.37 | 206.96 | 145.15 | 110.33 |
| 14      | CPD-14              | 261.96 | 215.17 | 186.33 | 155.11 |
| 15      | CPD-15              | 240.13 | 179.02 | 82.35  | 58     |
| 16      | CPD-16              | 253.02 | 203.31 | 163.26 | 115.27 |
| 17      | CPD-17              | 269.02 | 232.03 | 210.24 | 164.72 |
| 18      | CPD-18              | 236.04 | 171.04 | 172.26 | 49.4   |
| 19      | CPD-19              | 247.91 | 192.02 | 161.13 | 104.4  |
| 20      | Control             | 300   | 298.29 | 297.24 | 296.17 |
**Table 3a** Genome information of *Enterobacter Cloacae* Mr2

| Description                                      | MR2    |
|--------------------------------------------------|--------|
| DNA, total number of bases                       | 4758062|
| DNA coding number of bases                       | 4224707|
| DNA G+C number of bases                          | 2621395|
| Genes                                            | 5571   |
| Protein coding genes                             | 5404   |
| RNA genes                                        | 167    |
| rRNA genes                                       | 16     |
| 5S rRNA                                          | 8      |
| 16S rRNA                                         | 4      |
| 23S rRNA                                         | 4      |
| tRNA genes                                       | 72     |
| Protein coding genes with function prediction    | 4573   |
| without function prediction                      | 831    |
| Protein coding genes with enzymes                | 1115   |
| w/o enzymes but with candidate KO based enzymes  | 646    |
| Protein coding genes connected to Transporter Classification | 992    |
| Protein coding genes connected to KEGG pathways  | 1266   |
| not connected to KEGG pathways                   | 4138   |
| Protein coding genes connected to KEGG Orthology (KO) | 2423   |
| not connected to KEGG Orthology (KO)             | 2981   |
| Protein coding genes connected to MetaCyc pathways | 1085   |
| not connected to MetaCyc pathways                | 4319   |
| Protein coding genes with COGs                   | 2998   |
| in paralog clusters                              | 3861   |
| in Chromosomal Cassette                          | 5571   |
| Biosynthetic Clusters                            | 20     |
| Genes in Biosynthetic Clusters                   | 199    |
| Fused Protein coding genes                       | 99     |
| Protein coding genes coding signal peptides      | 446    |
| Protein coding genes coding transmembrane proteins | 1334   |
| COG clusters                                     | 1811   |
| KOG clusters                                     | 757    |
| Pfam clusters                                    | 2505   |
| TIGRfam clusters                                 | 1413   |
Table 3b COG categories of *Enterobacter cloacae* MR2 and *Enterobacter cloacae* subsp. cloacae ATCC 13047

| Class ID | Description | MR2 | ATCC13047 |
|----------|-------------|-----|-----------|
| A        | RNA processing and modification | 1   | 1         |
| W        | Extracellular structures | 2   | 4         |
| D        | Cell cycle control, cell division, chromosome partitioning | 43  | 50        |
| V        | Defense mechanisms | 88  | 59        |
| F        | Nucleotide transport and metabolism | 114 | 81        |
| Q        | Secondary metabolites biosynthesis, transport and catabolism | 145 | 114       |
| N        | Cell motility | 154 | 177       |
| I        | Lipid transport and metabolism | 162 | 131       |
| U        | Intracellular trafficking, secretion, and vesicular transport | 164 | 161       |
| O        | Posttranslational modification, protein turnover, chaperones | 175 | 158       |
| H        | Coenzyme transport and metabolism | 198 | 165       |
| L        | Replication, recombination and repair | 208 | 289       |
| J        | Translation, ribosomal structure and biogenesis | 227 | 206       |
| T        | Signal transduction mechanisms | 282 | 258       |
| M        | Cell wall/membrane/envelope biogenesis | 318 | 282       |
| C        | Energy production and conversion | 320 | 253       |
| S        | Function unknown | 390 | 383       |
| P        | Inorganic ion transport and metabolism | 431 | 351       |
| K        | Transcription | 470 | 437       |
| G        | Carbohydrate transport and metabolism | 596 | 463       |
| E        | Amino acid transport and metabolism | 657 | 500       |
| R        | General function prediction only | 718 | 626       |

Table 4 Assembly statistics genome sequence of *Enterobacter cloacae* MR2

| Sr. No. | Assembly Statistics | Value |
|---------|---------------------|-------|
| 1       | Total number of reads | 549,959 |
| 2       | Assembled Reads     | 507,383 |
| 3       | Coverage            | 18.06 X |
| 4       | Number of Contigs   | 230    |
| 5       | Consensus Length    | 4,758,062bp |
| 6       | Largest Contig      | 177,145bp |
| 7       | N50                 | 36,082bp |
| 8       | N90                 | 9,841bp |
| 9       | N95                 | 7,065bp |
**Table.5** RAST distribution of *Enterobacter cloacae* MR2 compare with *Enterobacter cloacae* subsp. cloacae ATCC 13047 and *Enterobacter* sp. 638

|                     | Enterobacter Cloacae |          |          |
|---------------------|----------------------|----------|----------|
|                     | MR2                  | ATCC 13047 | 638      |
| Cofactors, Vitamins, Prosthetic Groups, Pigments | 9.41 | 6.63 | 7.04 |
| Cell Wall and Capsule | 5.58 | 6.22 | 6.87 |
| Virulence, Disease and Defense | 2.85 | 3.71 | 2.64 |
| Potassium metabolism | 0.98 | 0.84 | 0.87 |
| Photosynthesis | 0.00 | 0.00 | 0.00 |
| Miscellaneous | 0.81 | 0.92 | 0.90 |
| Phages, Prophages, Transposable elements, Plasmids | 0.71 | 1.84 | 1.46 |
| Membrane Transport | 5.70 | 5.07 | 4.41 |
| Iron acquisition and metabolism | 2.02 | 2.23 | 1.40 |
| RNA Metabolism | 4.35 | 4.73 | 5.95 |
| Nucleosides and Nucleotides | 3.10 | 3.12 | 3.40 |
| Protein Metabolism | 6.29 | 6.32 | 8.02 |
| Cell Division and Cell Cycle | 0.69 | 0.72 | 1.07 |
| Motility and Chemotaxis | 2.93 | 3.89 | 2.61 |
| Regulation and Cell signaling | 3.83 | 3.61 | 3.84 |
| Secondary Metabolism | 0.12 | 0.13 | 0.11 |
| DNA Metabolism | 2.95 | 3.20 | 3.42 |
| Fatty Acids, Lipids, and Isoprenoids | 3.27 | 3.48 | 3.68 |
| Nitrogen Metabolism | 1.57 | 1.23 | 1.32 |
| Dormancy and Sporulation | 0.07 | 0.08 | 0.11 |
| Respiration | 3.98 | 3.97 | 3.96 |
| Stress Response | 4.50 | 4.89 | 4.97 |
| Metabolism of Aromatic Compounds | 1.35 | 0.97 | 0.14 |
| Amino Acids and Derivatives | 12.27 | 12.28 | 11.95 |
| Sulfur Metabolism | 1.60 | 1.59 | 1.85 |
| Phosphorus Metabolism | 1.30 | 1.20 | 1.46 |
| Carbohydrates | 17.75 | 17.17 | 16.55 |

**Table.6** Core and Pan genome analysis of *Enterobacter cloacae* MR2 with other *Enterobacter* genus spp

| Organism                  | CDS  | Pan CDS | Core CDS | Var CDS | Strain specific CDS |
|---------------------------|------|---------|----------|---------|---------------------|
| *E. cloacae* MR2          | 5703 | 5697    | 2089     | 3608    | 1876                |
| *Enterobacter* sp. 638    | 4396 | 4394    | 2056     | 2338    | 1062                |
| *E. cloacae* ENHKU01      | 4570 | 4566    | 2055     | 2511    | 681                 |
| *E. cloacae* subsp. dissolvens SP1 | 4682 | 4682    | 2052     | 2630    | 1848                |
| *E. cloacae* subsp. dissolvens SDM | 4758 | 4752    | 2055     | 2697    | 459                 |
| *E. cloacae* ATCC 13047   | 5707 | 5704    | 2061     | 3643    | 1311                |

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Fig.1 Per base sequence quality scores before (A) and after pre-processing and filtering (B)

Fig.2a Genes connected to the subsystems and their distribution in different categories

Annotation indicated 2907 features (genes or CDS) within 458 Subsystems and 4507 Coding Sequences

Fig.2b RAST distribution comparison Enterobacter cloacae MR2 with Enterobacter cloacae subsp cloacae ATCC 13047 and another neighbor genome Enterobacter sp. 638
**Fig. 3** Alignment of *Enterobacter cloacae* MR2 ordered contigs with the reference genome *Enterobacter cloacae subspp. cloacae* ATCC 13047

The alignment display is organized into one horizontal "panel" per input genome sequence. Each genome's panel contains the name of the genome sequence, a scale showing the sequence coordinates for that genome, and a single black horizontal center line. The regions of sequence with homology in the other two genome are indicated by colored blocks. The lines joining the blocks between three genomes trace each orthologous Locally Collinear Blocks (LCB) through every genome. In this case, Row1: *Enterobacter cloacae* MR2 ordered contigs, Row2: *Enterobacter cloacae subspp. cloacae* ATCC 13047 genome, Row3: *Enterobacter* spp. 638 genome.

**Fig. 4** *Enterobacter cloacae* MR2 graphical representation in MaGe’s genome browser and synteny maps
**Fig.5** Circular representation of the *Enterobacter cloacae* MR2 genome

Circles display (from the outside): (1) GC percent deviation (GC window - mean GC) in a 1000-bp window. (2) Predicted CDSs transcribed in the clockwise direction. (3) Predicted CDSs transcribed in the counterclockwise direction. Genes displayed in (2) and (3) are color-coded according different categories: red and blue: MaGe validated annotations, orange: MicroScope automatic annotation with a reference genome, purple: Primary/Automatic annotations. (4) GC skew (G+C/G-C) in a 1000-bp window. (5) rRNA (blue), tRNA (green), misc_RNA (orange), Transposable elements (pink) and pseudogenes (grey).

**Fig.6** Pan/Core Genome Analysis of *Enterobacter cloacae* MR2, its reference *Enterobacter cloacae* subsp *cloacae* ATCC 13047 and another neighbor genome *Enterobacter sp.* 638

Core-genome, variable-genome and strain specific sizes are represented with a Venn diagram. Values on diagram represent the numbers of MICFAM families for each organism intersections

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Clusters of orthologous genes and CDS have been highlighted in the map with the color intensity.

**Comparative genome analysis**

Comparative whole genome alignment is the key to find important regions to determine gene functions and establishing evolutionary events. Genome annotation identified plentiful of interesting genes that were involved in organo-phosphate degradation, detoxification and stress responses.

The BLAST based comparison of *E. cloacae* MR2 genome (Fig. 5) allowed sequence feature information to be visualized in context to reference sequence. Since this approach is reference based, only the regions present in the reference sequence or absent in the query sequence could be visualized. It also indicated that some of the features and sequences of the draft genome, which are missing from the reference genomes differed in significantly in terms of gene content. It shared most of the features in the reference genome, but there are regions and parts which might have deletions or missing of sequence parts.

Mauve software was used to construct genome-wide pair wise DNA alignments between *Enterobacter cloacae* MR2 and *Enterobacter* sp. 638. Synteny map was used for visualization of comparative analysis of complete genome assemblies at different levels of resolution, ranging from genome-scale comparison of chromosomes to comparisons of individual regions of alignment at the nucleotide level (Fig. 3).

The position of the blocks indicating the MR2 genome has different synteny as compare to other strains of the *Enterobacter*. The gap region between the block is also have the difference in the all three genomes.

**Pan/core genome analysis**

The Pan Genome analysis allowed to determine the common and variable genome proportion for each genome involve in the analysis. It also extracts core-genome, variable-genome and strain specific coding sequences. The Pan/Core genome analysis was carried out using the genome of *Enterobacter* sp. 638, *E. cloacae* subsp. cloacae ENHKU01, *E. cloacae* subsp. dissolvens SP1, *E. cloacae* subsp. dissolvens SDM, *E. cloacae* subsp. cloacae ATCC 13047 along with MR2. The core and pan genome interpretation is represented by Venn diagram which represents the numbers of genes/CDS in cluster of eight groups, out of which majority of the genes are shared (2764) among the three genomes (Fig. 6). MR2, ATCC 13047 and *Enterobacter* sp. 638 has 2089, 2061 and 2056 CDS respectively, for the core genome while MR2 has strain specific 1876 CDS compare to 1311 and 1062 CDS in ATCC 13047 and *Enterobacter* sp. 638 CDS (Table 6).

The Indian agriculture sector has been growing rapidly over the years. In order to control the insect from attacking their crops, pesticides are applied to lower the damages on crops and forestry products. Application of pesticides increases the cost of cultivation; on the other hand it greatly reduces the losses caused by pests and diseases, creating great economic benefits. However, pesticide residues can adversely affect ecosystems and human health and also cause serious environmental pollution.

Organophosphorus compounds cause short and long term environmental hazards and health problems. These pesticides affect the nervous system by disrupting the enzyme that regulates acetylcholine, a neurotransmitter. Organophosphorus pesticide is used to control a variety of sucking, chewing and boring
insects, spider mites, aphids and pests that attack crops like cotton, sugarcane, peanuts, tobacco, vegetables, fruits and ornamentals. Among the insecticides, monocrotophos, quinalphos and chlorpyrifos top the list of organophosphorus insecticides in the Indian market (Singh et al., 2003; Singh et al., 2004).

In this study, Nineteen isolates were initially screened which were found to be highly efficient in degrading chlorpyrifos and environmental stress tolerance. The mpd gene specific marker was amplified in all chlorpyrifos degrading isolates, except 2 isolates which indicated the presence of new or unidentified gene. The bacterial strains were tested for their ability to degrade chlorpyrifos by High Performance Liquid Chromatography (HPLC) and the results of these works have been published (Parmar et al., 2014). It was found that CPD-12 named as E. cloacae strain MR2 based on 16s rDNA analysis, was able to degrade up to 500 ppm of chlorpyrifos in 30 h. Exploring and analyzing the genome of E. cloacae strain MR2 have further paved our way for understanding the mechanisms of pesticide degradation and the genes, biochemical pathways, and metabolites involved in organophosphate degradation. RAST annotation provided us with the most closely related strains in respect to our bacterial strain which are E. cloacae ATCC 13047 and Enterobacter sp. 638. RAST distribution also indicates the highest subsystem distribution as compare to E. cloacae ATCC 13047 and Enterobacter sp. 638.

Mapping of MR2 with E. cloacae ATCC 13047 and Enterobacter sp. 638 mapped most of the contigs from our draft genome with the reference genome E. cloacae subsp. ATCC 13047. The regions mapped were variations in the location and had gap regions, indicating the genomic island's presence in the genome. The annotation also revealed the presence of genes for phosphorus metabolism, sulfur metabolism, metabolism of aromatic compounds, nitrogen metabolism, protein metabolism, potassium metabolism, iron acquisition and metabolism in the genome of strain MR2 which indicated the capability of the stain to promote plant growth and related activity.

The pan-genome describes the full complement of genes in a list of organisms. It is the union of all the gene families and specific genes of all the strains. It includes the core-genome containing gene families shared by all the organisms (intersection of gene families) and the variable-genome containing genes families shared by two or more organisms and strain specific genes. MR2 have higher strain specific genes as compared to E. cloacae ATCC 13047 and Enterobacter sp. 638. (Ren et al., 2010) characterized the genome of E. cloacae ATCC 13047 and showed that the chromosome carries seven operons involved in toxic heavy-metal resistance, including two sil operons, three ars operons, a mer operon, and a cop operon. MR2 genome when compared to the same through synteny, it indicated the presence of the same gene clusters genome.

The Enterobacter sp. 638 genome was sequenced and analyzed by Taghavi et al., (2010) and explained the plant growth promoting properties of the strain. While comparing genome of MR2 to sp.638, it also showed the presence of the gene clusters for the siderophores, Indol 3-acetic acid and phosphate solubilization. Hence the Enterobacter cloacae MR2 found during the course of this study, if used in the soil will not only degrade pesticide but will also increase the plant growth.

Nucleotide sequence accession numbers
This Whole Genome Shotgun project has been deposited at DDBJ / EMBL /GenBank under the accession ARYB0000000000. Bio-project registered under Accession: PRJNA203096 ID: 203096.

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