Bacterial Community Structure in the Asian Rice Gall Midge Reveals a Varied Microbiome Rich in Proteobacteria

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The Asian rice gall midge (ARGM) has emerged as a model gall forming pest of rice. The ARGM infestation of rice results in failure of panicle formation and economic loss. Understanding the molecular basis of ARGM-rice interactions is very crucial in order to control this devastating pest of rice. The current investigation was devised to identify bacterial communities present in the ARGM and in addition the bacterial diversity in the maggots during their interaction with susceptible or resistant rice varieties. Sequencing of 16S rRNA bacterial gene (V3-V4 region) revealed differences in the microflora of the ARGM maggots feeding on susceptible or resistant rice hosts. Results revealed that Wolbachia was the predominant bacterium in pupae and adults while Pseudomonas was predominant in maggots. Further, we observed that members of proteobacteria were predominant across all the samples. There was high species diversity in maggots isolated from susceptible rice and a high representation of unclassified bacteria in maggots isolated from resistant rice. This is the first study that reports variation of microbiome of the ARGM, based on host phenotype from which it was isolated, and results suggest that these variations could have an important role in host's susceptibility.

The Asian rice gall midge (ARGM) is one of the most destructive insect pests of rice1. As a member of the specialized gall making insect family Cecidomyiidae, it spends most part of its life cycle within the rice plant. Maggot hatching from egg is a tiny larva that twists and slides down the space between leaf sheaths facilitated by a thin film of water and moisture. Upon reaching the apical meristem or crown tissue, maggot initiates feeding by lacerating the tissue with its sternal spatula, secreting saliva and engulfing the oozing cell contents. This results in noticeable change in growth of surrounding tissue that encircles the maggot to form a gall chamber and form nutritive nurse cells (i.e. shifting of the metabolic process from a reproductive phase to a vegetative phase)1. Maggot molts twice and passes through three instars prior to pupation within the now tubular gall. Pupa is exceptionally active and moves upwards in the gall through wriggling movement, drills an exit hole at the tip of gall using its cephalic horns and partially protrudes out to facilitate adult emergence. Adult fly resembling mosquito with pink body has short life span of 1–2 days. Adults do not feed on rice but mate and lay eggs on plant within the short life span. Successful establishment of the insect in the plant and formation of gall renders that tiller sterile without further formation of panicles and grains that result in economic loss. However, several of the rice genotypes resist insect establishment and gall formation offering genetic host-plant resistance1. Gall flies lay eggs almost indiscreetly with reference to such genotypes and egg hatching or movement of hatched maggots into the plant to reach apical meristem is not hindered. However, maggot survival is dramatically reduced to nil within 48 h of feeding on resistant genotypes. Resistance in rice against the ARGM is often accompanied by tissue necrosis referred as hypersensitive response (HR+) and in few other cases where resistance is without accompanying necrosis (HR−)2. On the contrary, in a susceptible plant, maggots lacerate and feed at the meristematic region of the rice seedling resulting in formation of an enclosed tube like structure referred to as gall. Gall formation renders rice tillers sterile i.e. shifts the developmental process from a reproductive phase to a vegetative phase3.

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Various studies have described and defined the mechanism involved during rice defense and the ARGM counter defense but no study has reported the involvement of microbes in the ARGM-rice interaction. Moreover, insect associated microbiome is known to be involved in a three-way interaction between plant-insect-microbe and therefore, information regarding identity and diversity of bacterial community composition within the ARGM from both susceptible and resistant rice hosts is likely to contribute towards a better understanding of ARGM biology and gall midge-rice interactions as well. Here, we investigate and evaluate the ARGM-associated microbiome in an effort to understand this interaction and derive meaningful insights into this interaction to evolve novel strategies to control this important insect pest of rice.

We analysed the bacterial community composition in the Asian rice gall midge (ARGM) biotype 1 using a Next Generation Sequencing (NGS) protocol with Illumina-MiSeq platform with focus on the V3-V4 hypervariable region of 16S rRNA bacterial gene. We demonstrated the presence of specific bacterial community structure within each sample of the ARGM (isolated at different developmental stages and from resistant and susceptible rice host) and further showed that the bacterial community structure differed among the samples with reference to the developmental stages and the host. Furthermore, these observed variations of the microbiome of the ARGM, with reference to the host from which they were isolated, may have an influential impact on the nature of the interaction between ARGM and rice.

Results

PCR amplification of 16S V3-V4 hypervariable region from the Asian rice gall midge samples. A 460 bp fragment of the hypervariable V3-V4 region of the 16S rRNA was PCR amplified using specific universal primers (see Material and Methods section) from total genomic DNAs (OD260/280 ratio = 1.80–1.91 and OD260/230 ratio = 1.45–1.85) of the Asian rice gall midge biotype 1 (GMB1) adults (male and female), pupae and maggots isolated from TN1 and RP (Fig. 1a).

Analysis of Illumina-MiSeq raw reads. Libraries (of the V3-V4 region) prepared from the 460 bp PCR-amplified fragment (OD260/280 ratio = 1.75–1.92 and OD260/230 ratio = 1.50–1.82) from the different GMB1 samples (Supplementary Table S1) yielded total of 4.3 million raw reads on the Illumina-MiSeq sequencing platform.

Pre-processing of the raw reads for elimination of the primer containing sequences, possible adapter sequences, low quality reads and unique dual index barcodes sequences containing 5–7 nucleotides, yielded 4.1 million high quality reads. These high quality reads ranged from 94.0–96.1% across the GMB1 samples (Supplementary Table S2). Duplicate and chimeric reads were removed using CD-HIT-DUP tool to block over-estimated abundance of species, genera and or function. The non-overlapping reads were retained for further analysis (Supplementary Table S2).

Identification, assignment of operational taxonomic units (OTUs). All sequences from GMB1 samples were clustered into OTUs at 97% sequence similarity using QIIME (see ref. 43). A total of 7609 OTUs were identified that represented the bacterial species present in these samples (Table 1, Supplementary Data S1) at a confidence limit of 0.8 for assignment of each representative sequences. These assignments contained both specific bacterial species as well as non-assignable species that were labelled as unassigned OTUs. For all GMB1
samples, the rarefaction curves reached near plateau, indicating that the sampling depth and sequencing coverage were good (Fig. 1b). Maggotts from susceptible hosts (GMB1LS) showed highest number (2280) of OTUs followed by maggots from resistant hosts (GMB1LR) with 2161 OTUs, pupae (GMB1P) with 906 OTUs (Table 1, Supplementary Data S1). A heat map image of OTUs was generated using QIIME pipeline and results clearly illustrated the phylum-level microbial abundance and differences in the taxonomic composition among the GMB1 samples (GMB1F, GMB1M, GMB1L and GMB1LR) (Fig. 2). An interactive graphical overview of relative abundance of bacterial diversity and confidence limits within the complex hierarchies of OTUs classifications from phylum up to species level was obtained through the Krona chart (Supplementary Data S2).

**Diversity Index.** The diversity of microbial community structure in the GMB1 samples was estimated by alpha and beta diversity indices. We used 5 groups of alpha diversity (diversity within sample) indices to compare the species richness estimators (observed OTUs, Chao1 and PD_whole_tree) and diversity index (Shannon and Simpson index) of the microbiome from each GMB1 sample and these index values were significantly different between the samples (Table 1). Shannon information index of diversity ranged from 1.48 to 4.68, with an average of 3.04 and the observed Simpson index of diversity ranged from 0.54 to 0.84, with an average of 0.69 (Table 1). The Chao1 species richness estimator value revealed that OTUs richness varied from 1604 to 3247 with the highest value in GMB1LS among the GMB1 samples. In addition, the rarefaction curves also revealed the same pattern of the highest relative diversity being present in the GMB1LR. Significantly, more OTUs were observed in GMB1LS followed by GMB1LR (Table 1).

Beta diversity (using QIIME) analysis for assessing microbial diversity between the GMB1 samples (as represented by principal co-ordinate analysis (PCoA): unweighted and weighted UniFrac) was carried out. Unweighted and weighted UniFrac analysis accounted for the presence/absence of OTUs distribution between samples (GMB1LR, GMB1LS, GMB1M, GMB1F) along axes PC1, PC2, and PC3 (Fig. 3a). Bacterial variance in the GMB1 samples clearly separated GMB1LS, GMB1LR, GMB1P, GMB1F and GMB1M samples in unweighted (Fig. 3a) and weighted (Fig. 3b) plots. The first axis (PC1) showed maximum (40%) bacterial variation in GMB1LS and GMB1LR, followed by the second axis (PC2) 24% bacterial variation in GMB1P; and the third axis (PC3) 22% bacterial variation in GMB1F and GMB1M (Fig. 3a). And in 3-D weighted plot, more than 95% bacterial variation was found in GMB1LS and GMB1LR, followed by 3.71% bacterial variation in GMB1P and GMB1F and 0.37% bacterial variation in GMB1M (Fig. 3b). Similarly, Parallel unweighted and weighted UniFrac analysis revealed maximum bacterial variance in GMB1LS and GMB1LR samples respectively (Fig. 3c,d). PCoA weighted and unweighted UniFrac data was evaluated in pair-wise GMB1 samples comparisons (Tables 2 and 3) using Bonferroni corrected P-value. Bonferroni correction was based on randomization of sequence (100) assignments of each pair-wise GMB1 sample. Except one pair, GMB1F and GMB1M, Bonferroni corrected P-value 1.0 was calculated for weighted UniFrac analysis of all nine pair-wise GMB1 sample comparisons (Table 2) and Bonferroni corrected P-value for unweighted UniFrac analysis of all ten pair-wise comparisons were found to be P ≤ 1.0e-02 (Table 3).

**Dissimilarity matrix.** The dissimilarity between OTUs of the GMB1 samples ranged from 0.04–0.47. GMB1LS and GMB1LR showed highest dissimilarity between samples based on OTUs and the maximum similarity was between OTUs from adult female and pupa (dissimilarity index 0.04; Table 4).

**Bacterial distribution and its comparative study.** Different OTUs identified in GMB1 samples were categorised into different phyla, classes, orders, families, and genera (Table 5, Supplementary Data S3). The relative abundance of different bacterial phyla across all GMB1 samples were identified as follows: Proteobacteria, Actinobacteria, Firmicutes, Bacteroidetes, Acidobacteria, Chloroflexi, Cyanobacteria, TM7, Euryarchaeota, Thermi, NKB19 and Verrucomicrobia (Fig. 4a, Supplementary Data S3). The predominant phylum was Proteobacteria, accounting for 97.8% of the total bacterial sequences. This phylum alone contributed to 98.7%, 94.2%, 94.8%, 98.9%, and 99.1% of the total bacterial sequences in the samples GMB1F, GMB1LR, GMB1LS, GMB1M and GMB1P, respectively. Even though bacterial sequences representing Actinobacteria were second

| S. No. | Sample name | Shannon | Simpson | Chao1 | PD_whole_tree | Observed species |
|--------|-------------|---------|---------|-------|---------------|-----------------|
| 1.     | GMB1F       | 1.89    | 0.59    | 1604  | 51.92         | 906             |
| 2.     | GMB1M       | 2.51    | 0.67    | 1620  | 56.39         | 992             |
| 3.     | GMB1P       | 1.48    | 0.54    | 2321  | 82.78         | 1270            |
| 4.     | GMB1LS      | 4.63    | 0.84    | 3247  | 112.3         | 2280            |
| 5.     | GMB1LR      | 4.68    | 0.82    | 3140  | 115.3         | 2161            |
| Total  |             | 15.19   | 3.46    | 11932 | 418.69        | 7609            |

Table 1. Alpha-diversity indices of microbial community structure in the GMB1 samples. The diversity indices (Shannon and Simpson index) and species richness indices (Chao1, PD (phylogenetic diversity)_whole_tree and observed OTUs) were determined at 97% sequence similarity from each GMB1 sample. Maggot (GMB1LR and GMB1LS) samples showed highest alpha diversity among the GMB1 samples. GMB1F, rice gall midge biotype 1 female (adult); GMB1M, rice gall midge biotype 1 male (adult); GMB1P, rice gall midge biotype 1 pupae; GMB1LS, rice gall midge biotype 1 larvae isolated from susceptible host; GMB1LR, rice gall midge biotype 1 larvae isolated from resistant host.
most abundant sequences, it contributed approximately to 1.0% of the total bacterial sequences identified. The remaining sequences representing Firmicutes, Bacteroidetes, Chloroflexi, Planctomycetes, Acidobacteria, Verrucomicrobia, Cynobacteria, Fibrobacteres, Spirochaetes and others, together contributed to <1.0% of the total bacterial sequences (Supplementary Data S4). The major taxa, representing Proteobacteria in GMB1 samples (Fig. 4b), belonged to the genera *Wolbachia* (72.5%), *Pseudomonas* (9.8%), *Enterobacter* (1.9%), *Methylpila* (1.8%), unclassified bacteria (1.6%, belongs to f_Comamonadaceae), *Novosphingobium* (1.2%), unclassified bacteria (1.1%, belongs to f_Enterobacteraeaceae), *Acinetobacter* (0.9%), *Stenotrophomonas* (0.7%), unclassified bacteria (0.5%, belongs to f_Sphingomonadaceae), *Pantoea* (0.4%), *Variorax* (0.3%), unclassified bacteria (0.2%, belongs to f_Pseudomonadaceae), *Rheinheimera* (0.2%), *Breuvndimonas* (0.2%), *Agrobacterium* (0.2%), *Ramlilibacter* (0.2%), unclassified bacteria (0.2%, belongs to f_Oxalobacteraceae and o_Rhizobiales), *Delftia* (0.1%) and *Actinobacteria* (Corynebacterium (0.4%) and 0.1% of Propionibacterium, Kouria, Nesterenkonia and Citricoccus) (Supplementary Data S4).

In individual samples, *Wolbachia* sp. was predominant in GMB1F (89.8%), GMB1M (79.6%) and GMB1P (97.3%) while *Pseudomonas* sp. was the predominant in GMB1LR (20.9%) and GMB1LS (24.1%). Surprisingly, *Enterobacter* sp. was present at a higher level (7.2%) in GMB1M samples alone while in the other samples of GMB1 its presence was <1%. *Methylpila* sp. was present in higher levels in GMB1LS (8.1%) and GMB1LR (5.3%) while *Novosphingobium* sp. was observed in GMB1F (4.8%) and at 0.1% level in both GMB1LR and GMB1LS. *Acinetobacter* sp. was found at 1.6%, 1.4%, 1.3%, 0.5% and 0.4% levels in GMB1LS, GMB1LR, GMB1M, GMB1F and GMB1P samples, respectively. *Stenotrophomonas* sp. was found in GMB1M (2.5%), GMB1LS (0.2%) and GMB1LR (0.1%) (Supplementary Data S4). Approximately, 37% of the sequences across all samples were

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**Figure 2.** Heat map showing relative distribution and abundance of the OTUs among the GMB1 samples at phylum level.
unclassified due the unavailability of matching sequences in the GenBank and may represent unique unreported bacterial sequences (Table 6, Supplementary Data S4).

Analysis of abundant and rare species. The bacterial population present in GMB1 samples were represented by both abundant and rare species (the rare species were defined as those with a species frequency of <0.01% of the total population and the remaining ones were considered to be abundant species). In all the samples there was a preponderance of rare species except in GMB1LR. While the rare species were present at 48.2%, 51.7%, 65.3%, 72.1% and 80.1% in GMB1LR, GMB1LS, GMB1M, GMB1F and GMB1P, respectively, the abundant species were present at 19.8%, 27.9%, 34.6%, 48.3% and 51.7% in GMB1P, GMB1F, GMB1M, GMB1LS and GMB1LR, respectively (Fig. 4c).

A Venn diagram was plotted to calculate the number of specific and shared species among the GMB1 samples (Fig. 5). Across all the samples, 335 rare species and 123 abundant species were uniquely present while the rest (273 rare species and 186 abundant species) were shared among the samples (Fig. 5a–d, Supplementary Data S5).

Figure 3. Assessment of microbial diversity in the GMB1 sample using Principal Co-ordinate Analysis (PCoA) (a) unweighted 3-D visualization and (b) parallel visualization of UniFrac plots to reveal quantitative assessment of microbial diversity between the GMB1 samples. (c) Principal Co-ordinate Analysis (PCoA)-weighted 3-D visualization and (d) parallel visualization of UniFrac plots to reveal qualitative assessment of microbial diversity between the GMB1 samples.

Table 2. Pair-wise comparison of microbial diversity (based on Principal Co-ordinate Analysis (PCoA)-weighted UniFrac statistical test). The Bonferroni corrected P-value was calculated from 100 randomizations of bacterial OTUs assignments of the GMB1 samples.
Of the 186 abundant and 273 rare species shared between all the samples, 36 (which includes *Wolbachia* and *Pseudomonas*) of the abundant and 6 species (including *Acinetobacter*) of the rare were commonly present in all GMB1 samples.

**Dynamics of the presence of *Pseudomonas* and *Wolbachia* in developmental stages of the Asian rice gall midge.** To analyse the predominant genera in a developmental stage of the GMB1, we PCR amplified *Pseudomonas*-specific and *Wolbachia*-specific fragments using species-specific primers. The 618 bp *Pseudomonas*-specific fragment was observed in maggots (GMB1LS and GMB1LR) dissected 24 hours after infestation (hai). In contrast, we observed a weak amplification of the *Pseudomonas*-specific fragment from GMB1P, GMB1F and GMB1M (Fig. 6). Conversely, the 870 bp *Wolbachia*-specific fragment was predominant in GMB1P, GMB1F and GMB1M, while we observed less amplification of *Wolbachia*-specific fragment in maggots (GMB1LS and GMB1LR) (Fig. 6). However, it is yet to be verified whether this dynamic variability in the occurrence of *Pseudomonas* and *Wolbachia* during the different stages of the midge’s life cycle, hints at their probable importance during the developmental stages of the GMB1 within the host.

**Comparison of microbial communities between Asian rice gall midge and Hessian fly.** In an effort to understand and correlate the microbiome of the Asian rice gall midge microbial communities with that present in the Hessian fly (also a Cecidomyiid and a major pest of wheat), we carried out a comparative study using microbiome data of the Hessian fly in the public domain.

The identified OTUs of the first instar maggots of these two gall insects were compared (Table 7) at >97% similarity cutoff. Alpha-proteobacteria and gamma-proteobacteria were present at 78.7 and 16.1%, respectively in the Asian rice gall midge, while in the Hessian fly they were reported at 5.1 and 91.9%, respectively (similar trend observed in *Drosophila*). *Pseudomonas* sp. was present at 36.9% in the Asian rice gall midge and at 90.2% in the Hessian fly (Table 7).

**Discussion**

Currently, there is very little information with regard to the role of ARG microbiome in the gall midge-rice interaction. Microscopic first instar rice gall midge maggot and lack of artificial diets make the studies on ARGM
difficult. But more importantly, we expect that the identification of constituents of the microbiome of the rice gall midge would yield valuable information regarding their likely role in the gall midge’s interaction with its rice host, if any. In addition, it would be worth investigating whether gall midge maggots feeding on resistant rice host are pushed towards death due to a decrease in the number of beneficial microbes within them. Hence, the present study was initiated for the identification of microbial communities and to obtain a better understanding of the microbiome in the interaction between the rice and gall midge.

Our current study identified several distinct OTUs, based on the NGS sequencing of the V3-V4 region of the 16S rRNA gene, in 5 different samples of the ARGm. Members of Proteobacteria were the predominant bacterial constituents in all GMB1 samples and this could be due to the fact that during the feeding stage of the Asian rice gall midge, the maggots (GMB1LS and GMB1LR) could be actively recruiting Proteobacteria from the rice hosts and these then could be actively transferred from the maggots to the pupal stage and finally to adults. Moreover, GMB1LS and GMB1LR samples, representing the only feeding stage of the rice gall midge, showed the highest alpha diversity indices (Table 1). It may be noted that member of the Proteobacteria has been reported as the major bacterial population in the biosphere due to their efficient colonization ability. Proteobacteria have also been reported as the predominant bacteria in wild Drosophila—a polyphagous dipteran insect. This is true for hematophagous insects as well i.e. bed bugs and mosquitoes (Aedes albopictus and Aedes aegypti) as well as ticks.

Wolbachia sp and Pseudomonas sp formed the major constituents of the Proteobacteria in all the GMB1 samples analysed. Wolbachia sp has been described as a microbe with a capacity to change several traits of its host. However, it is not known whether Wolbachia, present within the gall midge, imparts any benefits to its host during the gall midge-rice interaction. It has been shown previously that phenotypes mediated by Wolbachia influence insulin signalling and glucose metabolism in Drosophila, male killing in ladybird (Adalia bipunctata) and butterflies (Acrea ecedon, Hypolimnas bolina), feminization of host in a wide variety of invertebrates including insects, spiders, crustaceans and nematode worms, sex ratio distortion and affect fecundity in terrestrial ispod Oniscus ascellus, manipulate sex life and boosts reproductive success in Ugandan butterflies. Also, an earlier study has shown that there is correlation between gall midge mitotypes and infection status of ARGm with respect to Wolbachia. However, currently there is no information whether Wolbachia present within the ARGm exerts any influence on the rice - gall midge interaction.

Pseudomonas, a prominent constituent of Proteobacteria, was also present in high numbers in all the GMB1 samples. Pseudomonas sp present in agricultural pests such as Plutella xylostella, Spodoptera exigua, Spodoptera litura, Trichoplusia ni is known to be involved in biopesticide breakdown. Therefore, it is likely to play a similar role in the Asian rice gall midge as well. Taking this observation further, we speculate that Pseudomonas may also play a role in the breakdown of plant defence biomolecules produced by the rice host in response to the gall midge infestation. However, this is yet to be verified. Furthermore, it has been shown that Pseudomonas play an important role in protecting silkworm (Bombyx mori) larvae from oxidative stress and bacterial infections. Whether a similar role is played by Pseudomonas in the rice gall midge is yet to be elucidated.

![Figure 4](https://www.nature.com/scientificreports/images/figure4.png)

**Figure 4.** Relative abundance of different bacterial communities among all the GMB1 samples at the level of (a) phyla and (b) genus. (c) Percentage of abundant and rare species among the different GMB1 samples.
The microbiome of the ARGM was made up of both rare and abundant species. The presence of these rare and abundant species which are unique to a particular sample, may be indicative of its crucial role in biological function of that particular stage in the life cycle of ARGM and might facilitate its interaction with the rice host. Maggots (GMB1LR) on resistant rice host survived only for 48–96 hours after infestation (hai) and during this time period maggots stop feeding and die and are thus unable to induce galls on such resistant rice. In contrast, maggots (GMB1LS) feeding on susceptible rice host continue feeding and are able to develop further, complete their life cycle and emerge from the galls as adults. Compared to the bacterial population observed in GMB1LS, the maggots (GMB1LR) dissected from resistant rice plant showed reduced bacterial population of those belonging to Actinobacteria, Bacteriodetes, Proteobacteria while there was an increase in the population of bacteria belonging to Cyanobacteria and Planctomycetes (Supplementary Table 3a,b, Supplementary Data S6). This reduction of several bacterial populations in GMB1LR within 48 hai could have a bearing on GMB1LR mortality as hypothesized for humans where host stability and demise are correlated to change in the indigenous microbiome34.

Insect diets are often nutrient-poor or incomplete and some insects have specialized cells, called bacteriocysts that house beneficial intracellular bacteria considered to be responsible for supplementing the otherwise nutrient-poor diets of insects35, 36. Facultative bacteria such as Enterobacter and lactic-acid bacteria have been found in GMB1 samples and both these have been reported to promote insect fitness by contributing essential amino acids and vitamins to their host. Further, during molting from 1st instar to 2nd instar and to subsequent instar, followed by pupa and finally into adults, associated proteobacteria could be involved in host survival by synthesizing and providing essential nutrients37 during the molting period which are probably essential during the non-feeding stages in the insect’s life cycle.

Insect metamorphosis is a metabolically dynamic and complex process and it is likely that the insect-associated microbes play a role in facilitating metamorphosis38. Moreover, it is believed that during the different stages of metamorphosis there is a sterilization process at each stage and insects acquire new set of microbes at each stage39, 40.

| S. No. | Distribution of unidentified OTUs into family (f)/order (o) GMB1 (%) | GMB1LR (%) | GMB1LS (%) | GMB1M (%) | GMB1P (%) |
|--------|---------------------------------------------------------------|-------------|-------------|------------|------------|
| 1.     | f_Comamonadaceae                                             | 0.0         | 6.3         | 6.3        | 0.0        | 0.0        |
| 2.     | f_Enterobacteriaceae                                         | 0.0         | 0.1         | 0.0        | 3.9        | 0.0        |
| 3.     | f_Caulobacteriaceae                                         | 0.0         | 4.5         | 4.0        | 0.0        | 0.0        |
| 4.     | f_Sphingomonadaceae                                         | 1.9         | 0.1         | 0.1        | 0.0        | 0.0        |
| 5.     | f_Pseudomonadaceae                                          | 0.1         | 0.2         | 0.2        | 0.6        | 0.0        |
| 6.     | f_Oxalobacteriaceae                                         | 0.0         | 0.5         | 0.7        | 0.0        | 0.0        |
| 7.     | o_Rhizobiales                                               | 0.0         | 0.6         | 0.7        | 0.0        | 0.0        |
| 8.     | f_Aeromonadaceae                                             | 0.2         | 0.2         | 0.3        | 0.0        | 0.1        |
| 9.     | f_Xanthomonadaceae                                           | 0.0         | 0.1         | 0.1        | 0.2        | 0.1        |
| 10.    | f_Morassellaceae                                             | 0.0         | 0.3         | 0.3        | 0.0        | 0.0        |
| 11.    | f_Alphaproteobacteria                                        | 0.3         | 0.0         | 0.0        | 0.0        | 0.0        |
| 12.    | f_Rhodobacteriaceae                                         | 0.0         | 0.2         | 0.2        | 0.0        | 0.0        |
| 13.    | f_Betaproteobacteria                                         | 0.0         | 0.2         | 0.2        | 0.0        | 0.0        |
| 14.    | f_Planococccaceae                                            | 0.0         | 0.1         | 0.1        | 0.0        | 0.0        |
| 15.    | f_Phylobacteriaceae                                          | 0.0         | 0.1         | 0.1        | 0.0        | 0.0        |
| 16.    | f_Alteromonadaceae                                           | 0.0         | 0.2         | 0.1        | 0.0        | 0.0        |
| 17.    | f_Rhodocyclaceae                                             | 0.0         | 0.2         | 0.1        | 0.0        | 0.0        |
| 18.    | f_Methylbacteriaceae                                         | 0.0         | 0.1         | 0.1        | 0.0        | 0.0        |
| 19.    | f_Rhizobiaceae                                               | 0.0         | 0.1         | 0.1        | 0.0        | 0.0        |
| 20.    | f_Micrococccaceae                                            | 0.0         | 0.1         | 0.1        | 0.0        | 0.0        |
| 21.    | f_Flavobacteriaceae                                          | 0.0         | 0.1         | 0.0        | 0.1        | 0.0        |
| 22.    | f_Ectothiorhodospira                                         | 0.0         | 0.1         | 0.1        | 0.0        | 0.0        |
| 23.    | f_Rhodospirillaceae                                         | 0.0         | 0.1         | 0.1        | 0.0        | 0.0        |
| 24.    | f_Erythrobacteriace                                           | 0.0         | 0.1         | 0.1        | 0.0        | 0.0        |
| 25.    | o_Myxooccales                                                | 0.0         | 0.1         | 0.1        | 0.0        | 0.0        |
| 26.    | f_Pirellulaceae                                              | 0.0         | 0.2         | 0.0        | 0.0        | 0.0        |
| 27.    | f_Bradyrhizobiace                                            | 0.0         | 0.1         | 0.1        | 0.0        | 0.0        |
| 28.    | f_Intrasporangiace                                           | 0.0         | 0.0         | 0.1        | 0.0        | 0.0        |
| 29.    | o_envOPS12                                                   | 0.0         | 0.0         | 0.1        | 0.0        | 0.0        |
| 30.    | f_A4b                                                        | 0.0         | 0.1         | 0.0        | 0.0        | 0.0        |
| 31.    | f_Nocardioidace                                              | 0.0         | 0.1         | 0.0        | 0.0        | 0.0        |
| 32.    | f_Chitinophagace                                             | 0.0         | 0.1         | 0.0        | 0.0        | 0.0        |
| 33.    | f_Cyclobacteriace                                            | 0.0         | 0.1         | 0.0        | 0.0        | 0.0        |
| TOTAL  | 2.5                                                          | 15.4        | 14.5        | 4.8        | 0.2        |

Table 6. Unclassified bacterial OTUs (upto 0.1% cut off) across the GMB1 samples.
Though the present study did not reveal metamorphic stage-specific microbes, results indicate fluctuations in the relative abundance of these microbes at each stage of life cycle of the midge. It is interesting to note that *Wolbachia* is predominant in GMB1F, GMB1M and GMB1P, *Pseudomonas* is predominant in GMB1LR and GM1LS and one reason for this observed phenomenon could be that the larval stages (GMB1LR and GM1LS) represent the only feeding stage in the life cycle of the Asian rice gall midge and therefore, *Pseudomonas* could be helping maggots in the breakdown of plant allelochemicals. In the subsequent non-feeding stages, the role of *Pseudomonas* may not be that predominant and therefore, not maintained. Moreover, data indicate that though *Wolbachia* is not predominant in GMB1LR and GM1LS samples, they are the second most abundant ones. Therefore, as the larval stages metamorphose to pupal and adult stages the population of *Pseudomonas* reduces and that of *Wolbachia* predominates in both the later stages. Certainly, concerted studies would be required to determine whether these microbes could be involved in bringing about changes in the internal morphology and physicochemical conditions that accompany metamorphosis in GMB1.

The results obtained from present study indicated that microbes present in the ARGM are predominantly members of the Proteobacteria. This is also true for another economically important member of Cecidomyiidae, the Hessian fly (*Mayetiola destructor*). It is also reasonable to assume that microbial diversity observed in GMB1...
is likely due to acquisition of different bacterial species from the external environment. The microbial diversity in GMB1 may influence GMB1-rice interaction and could contribute to gall formation in susceptible rice hosts. To the best of our knowledge, this is the first attempt of a comparative cataloguing of the microbiome in the different stages of the ARGM life cycle using a high-throughput next generation sequencing protocol. In addition, the extensive microbiome catalogue thus established (through data obtained from PCoA UniFrac analyses) will enable future studies involving association of the microbial genes with insect phenotypes and, even more broadly, its influence on insect interaction with its host and also metamorphosis.

**Methods**

**Sample Collection.** The GMB1 colony used in this study is being maintained in the greenhouse at the Indian Institute of Rice Research (IIRR) (formerly Directorate of Rice Research), Hyderabad, India under standard conditions. Briefly, pre-germinated seeds of different rice varieties were grown in rows, 5 cm apart, with 10–15 plants per row, in plastic trays with 8 cm deep, puddled, fertilizer-enriched, soil. 10–15-days-old plants were exposed to about 50 females and 10–20 males. On the third day, the trays were shifted to a high humidity (90–100% relative humidity) chamber and left undisturbed for egg incubation and maggot establishment. Maggots were isolated from the rice varieties TN1 and RP2068-18-3-5 (henceforth RP). GMB1 is virulent on TN1 (susceptible to GMB1; compatible interaction) and avirulent on RP (resistant to GMB1; incompatible interaction). Maggots feeding on TN1 complete their development and emerge as adults while those feeding on RP die within 96 h post hatching. Hence, first instar maggots were collected individually from both TN1 and RP within 96 h post hatching. Approximately 150 maggots from each rice variety (TN1 and RP) were collected using entomological needles and stored in 100% alcohol or RNAlater (Qiagen, USA) for further use. Care was taken not to injure the maggots. Approximately 150 maggots from each rice variety (TN1 and RP) were collected for the whole experiment. In addition, several pupae and adults were also collected from infested TN1. These stages will not be available, obviously, from RP.

**DNA extraction and PCR amplification of 16S ribosomal V3-V4 region from ARGM.** DNA was extracted from maggots, pupae and adults using HiPurA insect DNA purification kit (Himedia, India) as per manufacturer's instructions. DNA concentrations were estimated by NanoVue Plus (GE Healthcare, UK). 30 ng of extracted DNA was used as a template for PCR to amplify the V3-V4 region of the 16S rRNA region using the primers V3-341F 5′-CCTACGGGNGGCWGGCAG-3′ and V4-805R 5′-GACTACHVGGGTATCTAATCC-3′ (see Klindworth et al.42). PCR was carried out in 25 μl reaction volume containing 200 μM dNTPs, 5 U Taq DNA polymerase (Promega Corporation, USA) and 13 μM of each of the primers. PCR conditions were 95°C for 2 min, followed by 25 cycles at 95°C for 30 sec, 55°C for 30 sec, 72°C for 30 sec, with a final extension step at 72°C for 5 min. The amplified PCR products (~460 bp) from each sample were run on 1% agarose gel and gel-purified using QIAquick Gel Extraction kit (Qiagen, Inc, Germany) and quantified using NanoVue Plus spectrophotometer. The purified ~460 bp PCR products from each sample were sent to M/s Genotypic Technology Pvt. Ltd. (India) for metagenomic sequencing of the amplicon (16S rRNA V3-V4 region) using NGS protocol on an Illumina-MiSeq platform.

| S. No. | Reference          | Present study            | Bansal et al., 2014 |
|--------|--------------------|--------------------------|---------------------|
| 1.     | Insect             | Osorelia oryzae          | Mayetiola destructor |
| 2.     | Common name        | Asian rice gall midge    | Hessian Fly         |
| 3.     | Insect Order       | Diptera                  | Diptera             |
| 4.     | Member             | gall midge               | gall midge          |
| 5.     | Location           | Asia                     | USA                 |
| 6.     | Feed on            | Rice                     | Wheat               |
| 7.     | Platform used for microbial study | Illumina-MiSeq        | 454-Pyrosequencing |
| 8.     | Targeted 16S-rRNA hypervariable region | V3-V4                  | V3                  |
| 9.     | Raw Reads from 1st instar maggots | 782167                  | 6062                |
| 10.    | High quality reads | 739176                   | 3654                |
| 11.    | Developmental stages for microbial study | 1st instar maggots, pupa, adult male and female | P1, 2nd & 3rd instar larvae |
| 12.    | OTUs picked at     | >97% similarity cut-off  | >97% similarity cut-off |
| 13.    | Predominant proteobacteria in 1st instar | 97.6%                 | 96.33%              |
| 14.    | Alpha-proteobacteria in 1st instar | 78.7%                 | 5.08%               |
| 15.    | Beta-proteobacteria in 1st instar | 2.9%                   | 2.96%               |
| 16.    | Gamma-proteobacteria in 1st instar | 16.1%                 | 91.89%              |
| 17.    | Predominant bacteria in 1st instar | Pseudomonas sp., 36.9% | Pseudomonas sp., 90.2% |
| 18.    | Predominant bacteria in 2nd instar | ND                     | Buttiauxella sp., 9.7% |
| 19.    | Predominant bacteria in 3rd instar | ND                     | Buttiauxella sp.     |
| 20.    | Predominant bacteria in pupa | Wolbachia sp., 97.3% | ND                  |
| 21.    | Predominant bacteria in adult female | Wolbachia sp., 89.8% | ND                  |
| 22.    | Predominant bacteria in adult male | Wolbachia sp., 79.8% | ND                  |

**Table 7.** Comparative overview of microbial communities identified in maggots of the Asian rice gall midge and the Hessian fly. (ND; Not determined).

*Orseolia oryzae* and *Mayetiola destructor* are highly abundant in maggots of the Asian rice gall midge and the Hessian fly, respectively. This study provides a comparative catalogue of the microbiome of the Asian rice gall midge in different stages of its life cycle using next generation sequencing technology. This will serve as an essential platform for future studies aimed at understanding the role of microbiome in insect phenotypes and interactions.
Library Preparation and Illumina-MiSeq Sequencing. The library preparation followed the general library construction protocol as detailed in the 16S metagenomic sequencing library preparation methodology: (http://support.illumina.com/downloads/16s_metagenomic_sequencing_library_preparation.html). Gel purified fragment (~2 ng), of each sample was used for re-amplifying V3-V4 region of 16S rRNA region with specific primers that possess a tag sequence that are complementary to the Illumina sequence adapter and index primers from the Nextera XT Index kit V2 (Illumina, Cat # FC-131-2002) as per manufacturer’s protocol. This generated ~530 bp amplicons. Amplification was verified on 1% agarose gel before proceeding for indexing PCR. In the next round of PCR (indexing PCR), Illumina sequencing adapters and dual indexing barcodes were added using limited cycle PCR to generate a final product of ~600 bp. The library was cleaned using High Prep PCR clean up system (MagBio Genomics, Inc., USA). Further, quality control check of the library was performed using a Bioanalyzer Chip (Agilent Technologies, USA). PCR products with unique indices from each library were taken in equal (~2 ng) quantities and subjected to 300 paired-end multiplex sequencing using Illumina-MiSeq platform.

Initial Processing and bioinformatics analysis of sequence reads. QIIME (www.qiime.org)43 platform was used to extract high-quality sequences. Raw reads were selected based on read length, quality, primer and tag. The raw reads were quality checked using SeqQC v2.1 (genotypic.co.in/SeqQC.html), a proprietary software available with Genotypic Technology Pvt. Ltd, Bangalore, India. Based on the following criteria, the total sequences were screened to remove: (i) raw reads <110 nucleotides, (ii) reads with adapter sequences, (iii) raw reads with primer sequences, barcode and degenerate bases, (iv) reads that had a quality (phred) score of less than 20, (v) duplicates from processed reads using CD-HIT-DUP44, (vi) chimeric sequences from the operational taxonomic units (OTUs). The screened sequences were aligned to obtain representative sequences. The unique sequences set was classified into OTUs under threshold of 97% identity with phred score >20. The identity of each OTU sequence was confirmed using the microbial 16S ribosomal Greengenes database (Lawrence Berkeley National Laboratory, http://greengenes.lbl.gov). Next, a de novo taxonomic tree was constructed employing chimera-checked OTU set in QIIME (www.qiime.org)45,46 for downstream analysis which included estimation of alpha and beta diversity. The Shannon-Wiener47, Simpson’s diversity47, the Chao148 and rarefaction curve were calculated in order to evaluate the alpha diversity. Consequently, the phylogenetic relationships among the individuals were estimated to provide a complete picture of alpha diversity. UniFrac distance was based on the phylogenetic tree by QIIME. To estimate the dissimilarities between the different samples, principal co-ordinate analysis (PCoA) was carried out. Both weighted and unweighted calculations were performed for the PCoA (www.qiime.org).

Nucleotide Sequence accession number. Paired end Illumina sequence data from this study was submitted to the NCBI Sequence Read Archive (SRA) under accession number SRR064774.

PCR amplification of 16S ribosomal DNA (rDNA) of Wolbachia and Pseudomonas from the Asian rice gall midge samples. Wolbachia-specific primers29 (forward: 5′-TTGTAGCCGTAGTATGATACT-3′ and reverse: 5′-GAATAGTTAAGGATTTTACGT-3′) and Pseudomonas-specific primers29 (forward: 5′-GACCGGTGAGTAATGCCA-3′ and reverse: 5′-CAGCTGGTTGCTCTCCTATAA-3′) were used for PCR amplification of ~870 bp and ~618 bp 16S regions from Wolbachia and Pseudomonas, respectively to quantify their relative presence in the different developmental stages of the GMB1. The actin gene (~149 bp) of the ARGM 50 was used as the internal control. Genomic DNA (10 ng) of maggots, pupae and adults from TN1 and genomic DNA (20 ng) of maggots from RP were used for PCR. Twenty five μl PCR reaction mix contained 200 M dNTPs, 2.0 U of Taq DNA polymerase (Agilent Technologies, USA) and 12 μM of each primer. The 16S region of Wolbachia and actin gene were PCR amplified from each sample using the following conditions: an initial denaturation of 5 min at 95°C followed by 30 cycles of 30 s at 95°C, 1 min at 55°C, and 1 min at 72°C, with a final extension of 5 min at 72°C. The PCR amplified products were analysed on 1% agarose gel. The gels were photographed using a gel documentation system (Alpha Imager, Cell Biosciences, UK) and the images were captured in JPEG format. The images were subsequently analysed and relative intensities of each fragment was estimated using Image J software (http://imagej.nih.gov/ij/).

Statistical Analysis. Rarefaction analysis and alpha diversity index (Shannon and Simpson’s diversity index, Chao1) were used to estimate the differences in microbial communities between GMB1 samples (obtained from both TN1 and RP) used in the present study. The phylogeny-based unweighted UniFrac distance metric, and PCoA plots and rarefaction curves were plotted using QIIME (www.qiime.org). To visualize the relative abundance of bacterial phyla for individual GMB1 samples, heat maps were generated using QIIME.

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Author Contributions
S.N. designed the study. A.O. performed experiments and data analyses. A.O. and S.N. wrote the paper. J.S.B., D.K.S. and A.P.P. contributed reagents/materials and critically revised the manuscript. All authors have read and approved the final manuscript.

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