1	RAREFAN: a webservice to identify REPINs and RAYTs in bacterial genomes
2	Carsten Fortmann-Grote, Julia Balk and Frederic Bertels
3	
4	Max-Planck-Institute for Evolutionary Biology, Department of Microbial Population Biology
5	
6	
7	Corresponding author: Frederic Bertels, August-Thienemann-Straße 2, 24306 Plön, Germany,
8	bertels@evolbio.mpg.de.
9	
10	
11	
12	Running title: REPIN/RAYT Finder and ANalyzer
13	
14	Keywords: sequence analysis – mobile genetic elements – bacterial genomes –
15	Stenotrophomonas maltophila<u>maltophilia</u>
16	

17 Abstract

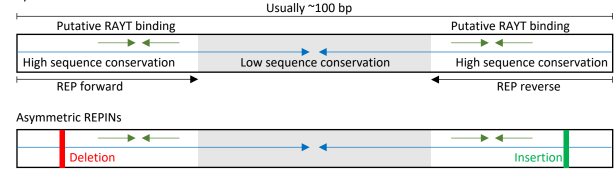
18 Compared to eukaryotes, mobile genetic elements repetitive sequences are rare in bacterial 19 genomes and usually do not persist for long-in the genome. Yet, there is at least one class of 20 persistent prokaryotic mobile genetic elements: REPINs. REPINs are non-autonomous 21 transposable elements replicated by single-copy transposases called RAYTs. REPIN-RAYT 22 systems are mostly vertically inherited and have persisted in individual bacterial lineages for 23 millions of years. Discovering and analyzing REPIN populations and their corresponding RAYT 24 transposases in bacterial species can be rather laborious, hampering progress in understanding 25 **REPIN-RAYT** biology and evolution. Here we present RAREFAN, a webservice that identifies 26 REPIN populations and their corresponding RAYT transposase in a given set of bacterial 27 genomes. We demonstrate RAREFAN's capabilities by analyzing a set of 49 Stenotrophomonas 28 maltophilia genomes, containing nine different REPIN-RAYT systems. We guide the reader 29 through the process of identifying and analyzing REPIN-RAYT systems across S. maltophilia, 30 highlighting erroneous associations between REPIN and RAYTs, and providing provide solutions 31 on how to find the correct associations. RAREFAN enables rapid, large-scale detection of 32 **REPINs and RAYTs, and provides insight into the fascinating world of intragenomic sequence** 33 populations in bacterial genomes.

35 Introduction

Repetitive sequences in bacteria are rare compared to most eukaryotic genomes. In eukaryotic
genomes, repetitive sequences are the result of the activities of persistent parasitic transposable
elements. In bacteria, in contrast, parasitic transposable elements cannot persist for long periods
of time (Park *et al.* 2021; van Dijk *et al.* 2022).(Park *et al.* 2021; van Dijk *et al.* 2022). To persist in
the gene pool, transposable elements musthave to constantly infect novel hosts (Sawyer *et al.*1987; Lawrence *et al.* 1992; Bichsel *et al.* 2010; Rankin *et al.* 2010; Wu *et al.* 2015; Park *et al.*2021). Yet, there is at least one exception: a class of transposable elements called REPINs.

43

Symmetric REPINs



44

45 Figure 1. The structure of symmetric and asymmetric REPINs. A typical REPIN consists of two 46 highly conserved regions at the 5' and 3' end of the REPIN (white), separated by a spacer region of lower sequence conservation (grey). The entire REPIN is a palindrome (blue arrows), which 47 means it can form hairpin structures in single stranded DNA or RNA. Each 5' and 3' region contains 48 49 a nested imperfect palindrome, which is referred to as REP (repetitive extragenic palindromic) sequence and has first been described in *Escherichia coli* (Higgins et al. 1982). REPINs can be 50 either symmetric or asymmetric. Asymmetric REPINs have a deletion and a corresponding 51 52 insertion in the highly conserved 5' or 3' end, which leads to "bubbles" in the hairpin structure. 53 REPINs in *E. coli* are asymmetric, which makes analyses with RAREFAN more challenging. REPINs (REP doublet forming hair PINs) are bacterial repetitive sequences that occur in extragenic spaces 54 55 (Bertels, Rainey 2011). REPINs are non-autonomous mobile genetic elements that are duplicated by a domesticated, single copy RAYT transposase (Nunvar et al. 2010; Bertels, Rainey 2011; Ton-56

Hoang *et al.* 2012). In contrast to typical bacterial mobile genetic elements, REPINs have persisted
for at least 100 million years in various species (Bertels, Gallie, *et al.* 2017; Park *et al.* 2021;
Bertels, Rainey 2022), in the absence of horizontal transfer of RAYT transposases or REPIN
populations (Bertels, Gallie, *et al.* 2017; Park *et al.* 2021; Bertels, Rainey 2022). These
evolutionary characteristics are consistent with REPIN-RAYT systems providing a benefit to the
host (Bertels, Rainey 2022).

The studyFigure adapted from (Bertels, Rainey 2022).

63

64 REPINs are short (~100 bp) nested palindromic sequences (Figure 1) that consist of two inverted 65 REP (repetitive extragenic palindromic (Higgins et al. 1982)) sequences that can be present 66 hundreds of times per genome (Bertels, Rainey 2011a). Most REPINs are symmetric where the 5' 67 REP sequences is identical to the 3' REP sequences, with the occasional substitution (Bertels, 68 Rainey 2011a; b). However, there are also asymmetric REPINs where the 5' REP sequence differs 69 from the 3' REP sequence by a point deletion or insertion (Bertels, Rainey 2011a, 2022), which makes the analysis and detection significantly more difficult (e.g., Escherichia coli REPINs). 70 Isolated REP sequences, REP singlets can also be found in the genome. These sequences are 71 72 decaying remnants of REPINs that are not mobile anymore (Bertels, Rainey 2011a). REPINs are 73 non-autonomous mobile genetic elements, which means they require a RAYT (REP Associated 74 tyrosine Transposase) transposase gene (also referred to as tnpAREP) to replicate inside the 75 genome (Nunvar et al. 2010; Bertels, Rainey 2011a; Ton-Hoang et al. 2012).

76

Within a genome, each REPIN population is usually only associated to a single RAYT gene. Hence,
 RAYT genes occur only in single copies per genome and do not copy themselves, unlike for
 example insertion sequences where often multiple identical sequences are present inside the
 genome. Unlike insertion sequences RAYT genes are also only inherited vertically, meaning they
 are host-beneficial transposases that are coopted by the host (Bertels, Gallie, *et al.* 2017; Bertels,
 Rainey 2022). The fact that REPINs and their corresponding RAYT genes are confined to a single

83	bacterial lineage makes them very special, in comparison to all other parasitic mobile genetic
84	elements in bacterial genomes (Bertels, Rainey 2022).
85	
86	Of a total of five different RAYT families, there are only two RAYT families that are associated
87	with REPINs: Group 2 and Group 3 RAYTs (Bertels, Gallie, et al. 2017). Group 2 RAYTs are present
88	in most Enterobacteria and usually occur only once per genome associated with a single REPIN
89	population. In contrast, Group 3 RAYTs are found in most Pseudomonas species and are usually
90	present in multiple divergent copies per genome, each copy associated with a specific REPIN
91	population (Bertels, Gallie, et al. 2017).
92	
93	REPINs and their corresponding RAYT genes occur exclusively in bacterial genomes and are
94	absent in eukaryotic or archaeal genomes (Bertels, Gallie, et al. 2017; Bertels, Rainey 2022).
95	Within bacterial genomes REPINs and RAYTs have been evolving in single bacterial lineages for
96	millions maybe even for a billion years (Bertels, Gallie, et al. 2017). The long term persistence of
97	REPINs in single bacterial lineages can also be observed when analyzing REPIN populations
98	(Bertels, Gokhale, et al. 2017; Bertels, Rainey 2022).
99	
100	Parasitic insertion sequences usually occur in identical copies in bacterial genomes reflecting the
101	fact that insertion sequences persist only briefly in the genome before they are eradicated from
102	the genome or kill their host (Park et al. 2021). REPINs in contrast are only conserved at the ends
103	of the sequence (presumably due to selection for function), the rest of the sequence is highly
104	variable and only the hairpin structure is conserved (Bertels, Rainey 2011a). The sequence
105	variability of REPINs within the same genome reflects their long-term persistence in single
106	bacterial lineages (Bertels, Rainey 2022). REPINs cannot simply reinfect another bacterial lineage
107	since they rely for mobility on their corresponding RAYT, which itself is immobile.
108	

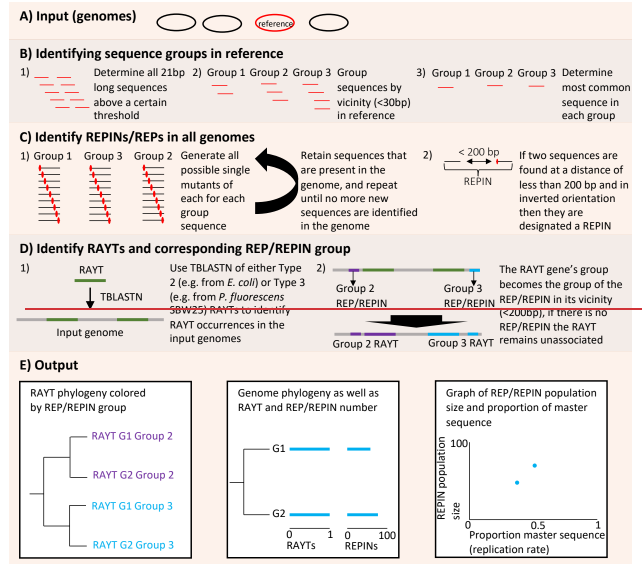
109 RAYTs and REPINs are distinct from typical parasitic insertion sequences, yet we know very little 110 about their evolution or biology. Currently, it is completely unclear what kind of beneficial 111 function maintains REPINs and RAYTs for millions of years in the genome. The reason for our lack 112 of knowledge is not because REPINs and RAYTs are rare. They are ubiquitously found in many 113 important and well-studied model bacteria such as Enterobacteria, Pseudomonads, Neisseriads, 114 Xanthomonads. Microbial molecular biologists presumably encounter REPINs quite frequently. 115 However, connecting the presence or absence of REPINs/RAYTs with phenotypes is difficult if we do not know when it is a REPIN that is present close to a gene of interest or a different type or 116 117 repeat sequence. Even if the scientist knows about the presence of a REPIN it is probably also 118 important to know whether a corresponding RAYT is present, since the function of REPINs largely 119 depends on the function of the presence of a corresponding RAYT gene (Bertels, Rainey 2022).

120

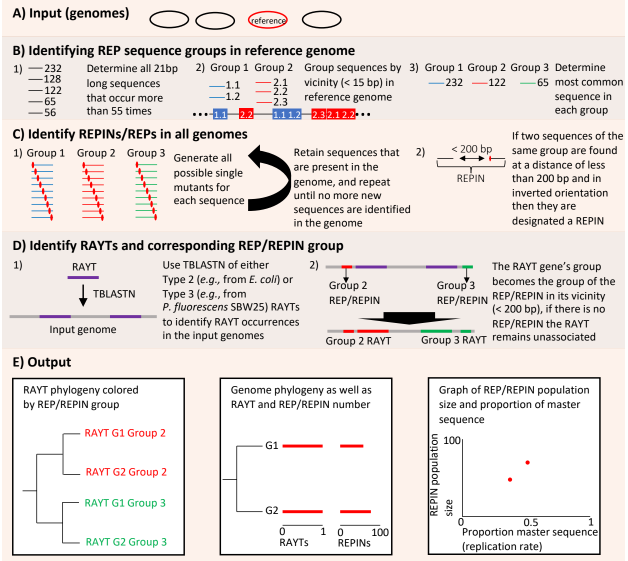
121 Yet, the identification of REPIN populations and their corresponding RAYTs can be rather 122 cumbersome. To facilitate REPIN studies, we have developed a webservice called if done from 123 scratch. This is particularly true if the microbial molecular biologist is not aware of all the ins and 124 outs of REPIN and RAYT biology. Identifying REPINs starts with an analysis of short repetitive 125 sequences in the genome. If there are excessively abundant short sequences present in the 126 genome, the distribution of these sequences is analyzed next. If they are exclusively identical 127 tandem repeats without sequence variation and present in only one or two loci in the genome 128 then these sequences are probably part of a CRISPR array and not REPINs.

129

Here, we present RAREFAN (RAYT/REPIN Finder and ANalyzer). Analyzer), a webservice that automates the identification of REPINs and their corresponding RAYTs. RAREFAN is publicly accessible at http://rarefan.evolbio.mpg.de and identifies REPIN populations and RAYTs inside <u>a</u> set of bacterial genomes. RAREFAN also generates graphs to visualize the population dynamics of REPINs, and assigns RAYT genes to their corresponding REPIN groups. Here we will demonstrate RAREFAN's functionality by analyzing REPIN-RAYT systems in the bacterial species *Stenotrophomonas maltophilia*.



And other files, for example: REP/REPIN and RAYT positions and sequences, REP/REPIN frequency, REP/REPIN conservation.



And other files, for example: REP/REPIN and RAYT positions and sequences, REP/REPIN frequency, REP/REPIN conservation.

Figure 1-2. RAREFAN workflow. (a) By default₇ RAREFAN requires the user to supply input sequences containing RAYTs and REPINs. These are, ideally, fully sequenced and complete genomes. (b) RAREFAN then identifies seed sequence groups (potential REP sequences) in the reference genome by first isolating all 21 bp (adjustable parameter) long sequences that occur more than 55 times (adjustable parameter) in the reference genome. It is likely that a large number of these sequences belong to the same REPIN sequence type, since the conserved part of REPINs is longer than 20bp. Hence, we grouped all sequences together that occur within 15 bp (adjustable parameter) of each other, anywhere in the genome. For example, if 'sequence 1' occurs 55 times and 'sequence 2' occurs 42 times then only one of these occurrences of 'sequence group. All further analyses are performed only with the most common sequence in each sequence group. This sequence will be called the seed sequence. (c) The occurrences of the seed and mutated seed sequences are identified in all submitted genomes. If a mutated seed sequence is identified in a genome, then all single mutants of that seed sequence are searched

recursively in the same genome. All identified sequences that occur within 130 bp in inverted orientation of each other are designated REPINs. All other identified seed sequences and mutated seed sequences are REP singlets. (d) TBLASTN is used to identify RAYT homologs (e-Value < 1e-30, adjustable parameter) of either E. coli (Group 2 RAYT) or from Pseudomonas fluorescens SBW25 (Group 3 RAYT) across all submitted genomes. If a RAYT homolog is in the vicinity (default 200 bp, adjustable parameter) of a previously identified REPIN or REP singlet, then this RAYT is designated as associated with this REPIN group. (e) Finally, RAREFAN plots three different summary graphs and(e) The first graph contains a RAYT phylogeny computed from a nucleotide alignment of all identified RAYT genes. The alignment is calculated with MUSCLE (Edgar 2004) and a phylogeny with PHYML (Guindon et al. 2010). The RAYT phylogeny indicates what RAYTs are associated with what REPIN populations (largest sequence cluster calculated with MCL) via colour coding. In a second graph the abundance of each REPIN population and RAYT copy number are displayed on a genome phylogeny. If no genome phylogeny is supplied RAREFAN calculates a whole genome phylogeny of the submitted genomes using andi (Haubold et al. 2015). In the last graph REPIN population sizes are plotted in relation to the proportion of master sequences. Master sequences are the most abundant REPIN in each population. The REPIN population is the largest sequence cluster that is formed by REPIN sequences (REP sequences are excluded). The largest sequence cluster is identified by applying MCL with an inflation parameter of 1.2 to a sequence matrix where only sequences are connected that differ in exactly one position (Van Dongen 2000). RAREFAN also generates various files containing, for example, REP, REPIN, or RAYT sequences and their positions in the query genomes.

137 Methods

138 Implementation

- 139 RAREFAN is a modular webservice. It consists of a web frontend written in the python 140 programming language (Van Rossum, Drake Jr 1995) using the flask framework (Grinberg 2018), 141 a java (Arnold et al. 2005) backend for genomic sequence analysis and an R (R Core Team 2016) 142 shiny app (RStudio, Inc 2013) for data visualization. The software is developed and tested under(R 143 Core Team 2016) shiny app (RStudio, Inc 2013) for data visualization. The software is developed 144 and tested on the Debian GNU/Linux operating system (Kleinmann et al. 2021). All components 145 are released under the MIT opensource license (Initiative 2021) and can be obtained from our 146 public GitHub https://github.com/mpievolbiorepository at
- 147 <u>scicom/rarefanhttps://github.com/mpievolbio</u>
- 148 <u>scicomp/rarefan.https://github.com/mpievolbio-scicomp/rarefan.</u>

- 149 The public RAREFAN instance at http://rarefan.evolbio.mpg.de runs on a virtual cloud server with
- 150 <u>4 single-threaded CPUs and 16GB of shared memory provided and maintained by the Gesellschaft</u>
- für Wissenschaftliche Datenverarbeitung Göttingen (GWDG) and running the Debian GNU/Linux
- 152 Operating System (Kleinmann *et al.* 2021).
- 153 The java backend drives the sequence analysis. It makes system calls to TBLASTN (Altschul *et al.*
- 154 1990) to identify RAYT homologs and to MCL (Van Dongen 2000) for clustering REPIN sequences-
- 155 <u>in order to determine REPIN populations.</u>
- Jobs submitted through the web server are queued and executed as soon as the required resources become available. Users are informed about the status of their jobs. After job completion, the user can trigger the R shiny app to visualize the results.
- 159 The java backend can also be run locally *via* the command line interface (available for download
- 160 at <u>https://github.com/mpievolbio-scicomp/rarefan/releases</u>).
- 161 Usage of the webservice
- The front page of our webservice allows users to upload their bacterial genomes in FASTA (.fas) format (**Figure 1A2A**). Optionally, users may also provide RAYT protein FASTA sequences (.faa) or phylogenies in NEWICK (.nwk) format. After successful completion of the upload process, the user fills out a web form to specify the parameters of the algorithm:
- Reference sequence: Which of the uploaded genome sequences will be designated as
 reference genome (see below for explanations). Defaults to the first uploaded filename
 in alphabetical order.
- Query RAYT: The RAYT gene that is used to identify homologous RAYTs in the query
 genomes.
- Tree file: A phylogenetic tree of the reference genomes that can be provided by the user,
 otherwise the tree will be calculated using andi (Haubold *et al.* 2015).
- Seed sequence length: The seed sequence length (in base pairs) is used to identify REPIN
 candidates from the input genomes. Default is 21 bp.

- Minimum seed sequence frequency: Lower limit on seed sequence frequency in the
 reference genome to be considered as a REP candidate. Default is 55.
- Seed sequence length: The seed sequence length (in base pairs) is used to identify REPIN
 candidates from the input genomes. Default is 21 bp.
- Distance group seeds: The maximum distance between a single occurrence of short
 repetitive sequences to still be sorted into the same sequence group.
- Association distance REPIN-RAYT: The maximum distance at which a REP sequence can
 be located from a RAYT gene to be linked to that RAYT gene.
- e-value cut-off: Alignment e-value cut-off for identifying RAYT homologs with TBLASTN.
 Default is 1e-30.
- Analyse REPINs: Ticked REPINs will be analysed (two inverted REP sequences found at a distance of less than 130 bp), if not ticked only short repetitive 21 bp long sequence will be analysed.
- User email (optional): If provided, then the user will be notified by email upon run
 completion.

The job is then ready for submission to the job queue. Upon job completion, links to browse and to download the results, as well as a link to a visualization dashboard are provided. If a job runs for a long time then users may also come back to RAREFAN at a later time, query their job status and eventually retrieve their results by entering the run ID into the search field at <u>http://rarefan.evolbio.mpg.de/results</u>. Relevant links and the run ID are communicated either on the status site or by email if the user provided their email address during run configuration. <u>Runs</u> <u>are automatically deleted from the server after six months.</u>

197 Identification of REPs and REPINs

The algorithm to determine REP sequence groups has been described in previous papers and is slightly improved in our implementation (Bertels, Rainey 2011, 2022; Bertels, Gokhale, *et al.* 200 2017). First, all N bp (21 bp by default) long seed sequences that occur more than M times (55 by default) are extracted from the reference genome. N and M are the seed sequence length and minimum seed sequence frequency, respectively (Figure 1B). All sequences occurring within the
 reference genome at least once within 15 bp of each other are then grouped together into n REP
 sequence groups (numbered 0 (n 1)). The most common sequence in each group, named REP
 seed sequence, is used for further analyses in each input genome.

206 In the next step all possible point mutants of the seed sequences are generated and searched for

207 in the genome (Figure 1C). The algorithm to determine REP sequence groups has been described

208 in previous papers and is slightly improved (Bertels, Rainey 2011a, 2022; Bertels, Gokhale, et al.

209 <u>2017). In our current implementation REPs/REPIN populations are now automatically linked to</u>

210 <u>RAYT genes.</u>

211 First, all N bp (21 bp by default) long seed sequences that occur more than M times (55 by default)

are extracted from the reference genome. N and M are the seed sequence length and minimum

seed sequence frequency, respectively (Figure 2B). All sequences occurring within the reference
 genome at least once within 15 bp of each other are then grouped together into n REP sequence

groups (numbered 0-(n-1)). The most common sequence in each group, named REP seed
 sequence, is used for further analyses in each input genome.

In the next step all possible point mutants of the seed sequences are generated and searched for
 in the genome (Figure 2C). If a sequence is found in the genome, then all possible point mutations
 are generated for this sequence as well and so on until no more sequences can be identified. If
 two sequences are found within 130 bp of each other in inverted orientation, then these are
 designated REPINS.

222 Identification of RAYTs

RAYTs are identified using TBLASTN (Camacho *et al.* 2009) with either a protein sequence
provided by the user or a Group 2 RAYT from *Eschericha coli* (yafM, Uniprot accession Q47152)
or a Group 3 RAYT from *P. fluorescens* SBW25 (yafM, Uniprot accession C3JZZ6). The presence of
RAYTs in the vicinity of a particular REPIN can be used to establish the association between the
RAYT gene and a REPIN group (Figure 1D).

228 <u>Among all identified REP and REPIN sequences REPIN populations can be isolated. REPIN</u>

229 populations are determined by applying MCL using an inflation parameter of 1.2 (Van Dongen

2000) to a network of REP/REPIN sequences where all sequences that differ by exactly one
 nucleotide are connected. The clustering results are stored in a file ending in .mcl. The
 sequences of the largest REPIN population (excluding REP singlets) are isolated in a file ending in
 largestCluster.nodes. The largest REPIN populations are shown in the REPIN population
 plot and the master sequence correlation plot (Figure 4).

235

236 Identification of RAYTs

237 RAYTs are identified using TBLASTN (Camacho et al. 2009) with either a protein sequence 238 provided by the user, a Group 2 RAYT from E. coli (yafM, Uniprot accession Q47152) or a Group 239 3 RAYT from *P. fluorescens* SBW25 (yafM, Uniprot accession C3JZZ6). The presence of RAYTs in 240 the vicinity (default 200 bp) of a particular REPIN can be used to establish the association 241 between the RAYT gene and a REPIN group (Figure 2D). All positions of all REPINs and REP 242 sequences of a REPIN group are checked whether they occur within 200 bp (by default) of a RAYT 243 gene. If so then the RAYT gene is linked to the REPIN group in the file 244 repin rayt association.txt.

245 Visualizations

For each REPIN-RAYT group summary plots are generated. These include plots showing the RAYT phylogeny, REPIN population sizes in relation to the genome phylogeny (calculated from a nucleotide alignment using MUSCLE (Edgar 2004) and PHYML (Guindon *et al.* 2010) to generate a phylogeny), REPIN population sizes in relation to the genome phylogeny (provided by the user or if not provided calculated by andi (Haubold *et al.* 2015)) as well as the proportion of master sequences (most common REPIN in a REPIN population) in relation to REPIN population size (Figure 1E2E).

253 Other outputs

Identified REPINs, REP singlets as well as RAYTs are written to FASTA formatted sequence files and to tab formatted annotation files that can be read with the Artemis genome browser (Rutherford *et al.* 2000). The REPIN-RAYT associations as well as the number of RAYT copies per genome are written to tabular data files. A detailed description of all output files is provided in the manual (<u>http://rarefan.evolbio.mpg.de/manual</u>).) and in the file "readme.md" in the output
<u>directory.</u>

260

261 Sequence analysis and annotation

262 For verification of RAREFAN results, REPIN-RAYT-systems were analyzedanalysed in their 263 corresponding genomes using Geneious Prime (Java Version 11.0.12+7 (64 bit); 264 Biomatters).version 2022.2.2 (Kearse et al. 2012). Nucleotide sequences and positions of REP 265 singlets, REPINs, and RAYTs were extracted from output files generated by RAREFAN and mapped 266 in the relevant S. maltophilia genome. The association of a RAYT gene to a REPIN population has 267 been validated when the corresponding seed sequence is flanking both ends of the RAYT gene 268 within 130 bp. Complete RAREFAN data used for analysis can be accessed by using the run IDs 269 listed in Table 1.

270

Table 1.- RAREFAN IDs linking to the raw data of the presented analyses.

Run ID	Reference genome
_1a8l7wu	S. maltophilia Sm53
<u>mknhxpv8</u>	S. maltophilia AA1
pgfmaxx5	S. maltophilia FDAARGO_649
<u>yy72i755</u>	S. maltophilia AB550
<u>78eu9zl0</u>	S. maltophilia ISMMS3

Associated data can accessed by entering the run ID at <u>http://rarefan.evolbio.mpg.de/results</u>.

273

274 Results

RAREFAN can identify REPINs and their corresponding RAYTs in a set of <u>- ideally</u> fully sequenced
 -bacterial genomes. The RAREFAN algorithm has been used in previous analyses to identify and
 characterize REPINs and RAYTs in *Pseudomonas* (Bertels, Rainey 2011, 2022), *Neisseria* (Bert

277 Characterize REFINS and RATTS III - Seducitionus (berteis, Rainey 2011, 2022), Neissena (berteis,

278 Rainey 2022), and Enterobacteria (Bertels, Gallie, et al. 2017; Park et al. 2021)Pseudomonads

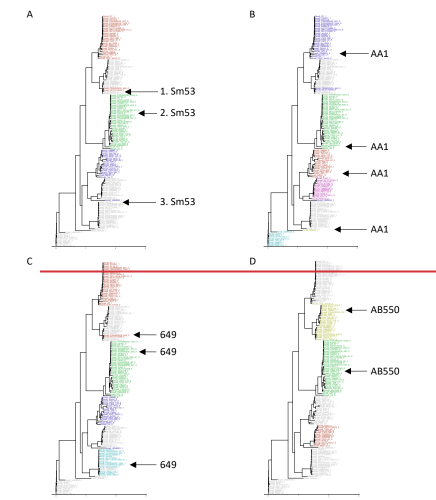
279 (Bertels, Rainey 2011a, 2022), Neisseriads (Bertels, Rainey 2022), and Enterobacteria (Bertels,

280 <u>Gallie, et al. 2017; Park et al. 2021</u>. To demonstrate RAREFAN's capabilities, we are presenting

an analysis of 49 strains belonging to the opportunistic pathogen *S. maltophilia*.

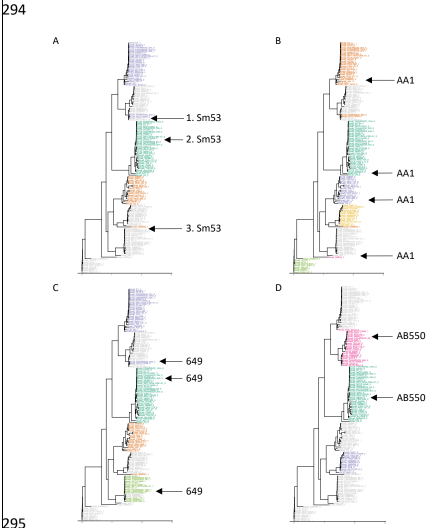
S. maltophilia strains contain Group 3 RAYTs, which are also commonly found in plant associated
 Pseudomonas species such as *P. fluorescens* or *P. syringae* (Bertels, Rainey 2011, 2022). Similar
 to Group 3 RAYTs in other species, *S. maltophilia* contains multiple REPIN RAYT systems per
 genome. Group 2 RAYTs, in contrast, contain only ever one REPIN RAYT system per genome
 (Bertels, Rainey 2022).

287



288

S. maltophilia strains contain Group 3 RAYTs, which are also commonly found in plant-associated
 Pseudomonas species such as *P. fluorescens* or *P. syringae* (Bertels, Rainey 2011a, 2022). Similar
 to Group 3 RAYTs in other species, *S. maltophilia* contains multiple REPIN-RAYT systems per
 genome. Group 2 RAYTs, in contrast, contain only ever one REPIN-RAYT system per genome
 (Bertels, Rainey 2022).



295

Figure 23. Phylogenetic trees built from RAYT genes extracted from *S. maltophilia* genomes. RAYT genes are coloured according to their association with REPIN populations in the reference genome. If noa REPIN population of a query genome is not present in the reference genome is associated with a, then the REPIN population cannot be identified in the query genome and the corresponding RAYT gene the RAYT genecannot be linked and is coloured in grey. The four panels A-D show phylogenies for four different reference strains. S. maltophilia strains Sm53, AA1, 649 and AB550 were used in panels A to D, respectively. Locations of a reference strain's RAYT genes in the tree are indicated by arrows. An association between almost all RAYTs and REPIN populations could be made by using four different reference genomes. Most of the RAYT genes are coloured (associated to a REPIN group) in at least one of the trees. The three numbered RAYT genes from the Sm53 RAREFAN run are referenced in the text.

297 Nine different REPIN-RAYT systems in S. maltophilia

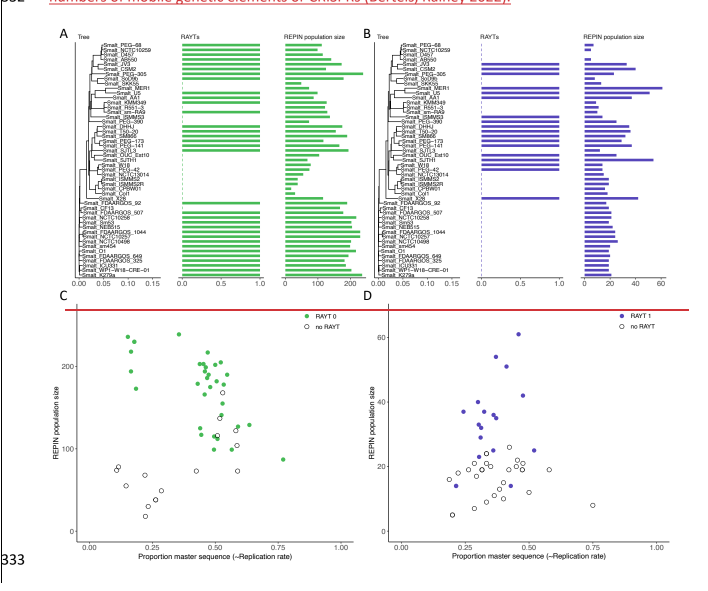
REPIN-RAYT systems in *S. maltophilia* are surprisingly diverse compared to other species. For example, *P.Pseudomonas* chlororaphis contains three separate REPIN populations that are present in all *P. chlororaphis* strains, each associated with its cognate RAYT gene (Bertels, Rainey 2022). *S.*(Bertels, Rainey 2022). *S. maltophilia*, in contrast, contains only one REPIN-RAYT system that is present across almost the entire species (green clade in Figure 23), and at least eight REPIN-RAYT systems that are present in subsets of strains (nine clades in Figure 45).

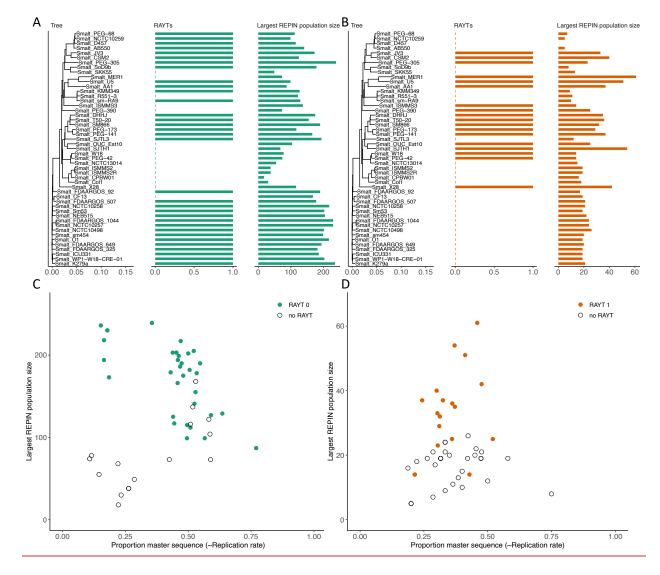
304 The patchy presence-absence pattern of REPIN-RAYT systems in *S. maltophilia*, makes the dataset 305 quite challenging to analyse. If a REPIN population is not present in the reference strain then 306 RAREFAN will not be able to detect it in any other strain. Yet, it is possible to detect RAYT genes 307 in all strains of a species independent of the reference strain selection. RAYT genes that are not 308 associated to a REPIN population are displayed in grey (Figure 2A3A). While these RAYT genes 309 are not associated to REPIN populations detected in the reference strain, they might still be 310 associated with a yet unidentified REPIN type- present in the genome the unassociated RAYT 311 gene is located in.

In order to identify all REPIN populations across a species, we suggest to perform multiple
 RAREFAN runs with different reference strains. The RAREFAN web interface supports re launching a given job with modified parameters.

³¹⁵ _To identify as many different REPIN-RAYT systems as possible in each subsequent run the ³¹⁶ reference should be set to a genome that contains RAYTs that were not associated with a REPIN ³¹⁷ population previously (*i.e.*, genomes containing grey RAYTs in **Figure 23**). However, this strategy ³¹⁸ may also fail when the REPIN population size falls below the RAREFAN seed sequence frequency ³¹⁹ threshold.

For example, *S. maltophilia* Sm53 contains three RAYTs only one of which is associated with a REPIN population (RAYT genes indicated in **Figure 2A3A**). However, the remaining two RAYTs are indeed associated with a REPIN population, but these REPIN populations are too small to be detected in *S. maltophilia* Sm53 (the seed sequence frequency threshold is set to 55 by default). In other *S. maltophilia* strains the REPIN populations are large enough to exceed the threshold. 325 For example, if *S. maltophilia* AB550 is set as reference, RAYT number 1 from Sm53 (Figure 2A3A) 326 is associated with the yellowpink REPIN population (Figure 2D3D). If S. maltophilia 649 is set as 327 reference RAYT number 3 from Sm53 (Figure 2A3A) is associated with the turquoiselight green 328 REPIN population (Figure 2C3C). RAYTs from the bottom clade are only associated with REPIN 329 populations when S. maltophilia AA1 is chosen as reference (Figure 2B).3B). While lower 330 thresholds can guarantee that all REPINs will be identified in the genome, the number of 331 sequence groups that are not REPINs quickly explodes. Especially for genomes that contain large numbers of mobile genetic elements or CRISPRs (Bertels, Rainey 2022). 332





334

Figure 34. REPIN population sizes and conservation. The plots show two REPIN populations and their associated RAYTs that were identified in *S. maltophilia* using *S. maltophilia* Sm53 as reference. (**A**) The phylogenetic tree on the left side is a whole genome phylogeny generated by andi (Haubold *et al.* 2015). Shown on the right are REPIN population sizes (which is the largest <u>REPIN cluster calculated by MCL)</u> and the number of associated RAYTs sorted by the genome phylogeny. The green REPIN populations and associated RAYTs are present in most strains in high abundance (maximum of 239 occurrences in *S. maltophilia* K279a, left panel). (**B**) The blueorange population in contrast is present in much lower numbers (maximum of 61 occurrences in *S. maltophilia* MER1, right panel). Note, REPIN populations are assigned consistent colours based on their abundance in the reference genome. For example, the most abundant REPIN population is always coloured in green, and the second most abundant population is always coloured in green, and the second most abundant population is always coloured in a REPIN population is the most common REPIN sequence. The master sequence in a REPIN population is the most common REPIN sequence. Thele an equilibrium the higher the proportion of the master sequence in the population the higher the replication rate (Bertels, Gokhale, *et al.* 2017). The presence and absence of an

associated RAYT is also indicated by the colours of the dots. Empty circles indicate that the REPIN population is not associated with a RAYT gene in that genome.

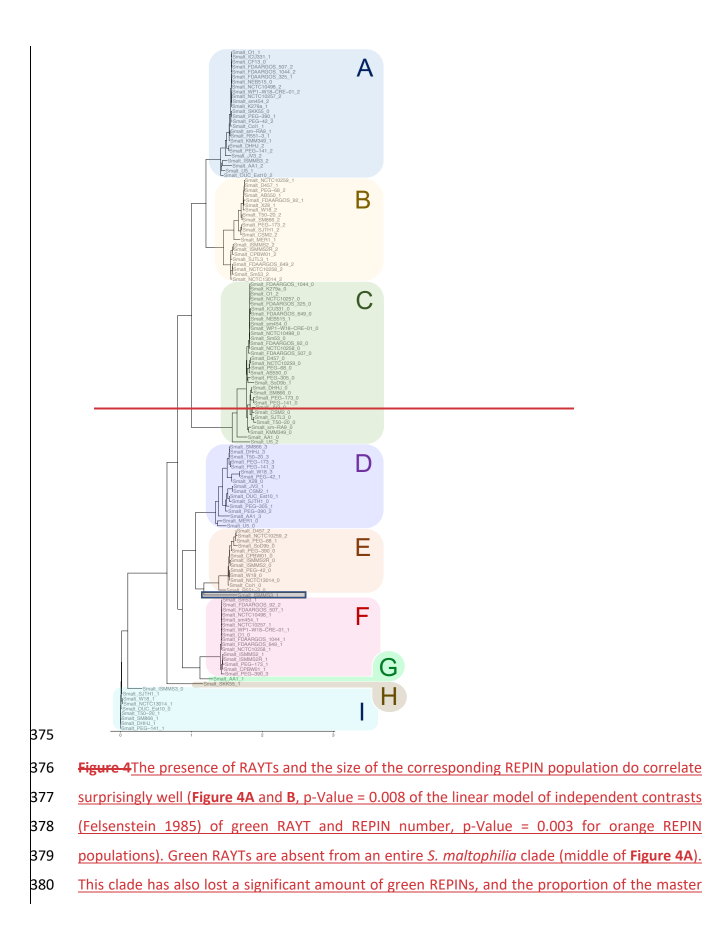
B35 RAREFAN visualizes REPIN population size and <u>potential</u> replication rate

336 The RAREFAN webserver visualizes REPIN population size and RAYT numbers in barplots. The 337 barplot is Barplots are ordered by the phylogenetic relationship of the submitted bacterial strains 338 (Yu et al. 2018). RAREFAN detects three populations when S. maltophilia Sm53 is selected as 339 reference strain (Figure 2A3A). The largest REPIN population (calculated by MCL from all REPINs 340 of that type) has a corresponding RAYT gene in almost all strains (first barplot in Figure 3A4A) 341 and most REPIN populations contain more than 100 REPINs (second barplot in Figure 3A4A). The 342 second largest REPIN population in Sm53 (purpleorange population in Figure 3B4B) is 343 significantly smaller and contains no more than 61 REPINs in any strain and most strains do not 344 contain a corresponding RAYT for this population.

345 RAREFAN also provides information on REPIN replication rate (Figure 3-C4C and D). REPIN 346 replication rate can be estimated by dividing the number of the most common REPIN sequence 347 (master sequence) by the REPIN population size if the population is in mutation selection balance 348 (Bertels, Gokhale, et al. 2017). If a REPIN replicates very fast most of the population will consist 349 of identical sequences because mutations do not have enough time to accumulate between 350 replication events. Hence, the proportion of master sequences will be high in populations that 351 have a high replication rate. Transposable elements such as insertion sequences consist almost 352 exclusively of identical master sequences because the time between replication events is not 353 sufficient to accumulate mutations and because quick extinction of the element usually prevents 354 the accumulation of mutations after replication (Park et al. 2021; Bertels, Rainey 2022)(Park et 355 al. 2021; Bertels, Rainey 2022). REPIN populations in contrast replicate slowly and persist for long 356 periods of time, which means that a high proportion of master sequences suggests a high REPIN 357 replication rate.

In *S. maltophilia* the proportion of master sequences in the population does not seem to correlate well with REPIN population size, both in the green and the <u>purpleorange</u> population (**Figures 360 3C4C and D**). Similar observations have been made in *P. chlororaphis* (Bertels, Rainey **361 2022**)(Bertels, Rainey 2022), and may suggest that an increase in population size is not caused by an increase in replication rate. Population size is likely to be more strongly affected by other factors such as the loss of the corresponding RAYT gene, which leads to the decay of the REPIN population. One could even speculate that high REPIN replication rates are more likely to lead to the eventual demise of the population due to the negative fitness effect on the host (Bertels, Rainey 2022)(Bertels, Rainey 2022).

367 The presence of RAYTs and the size of the corresponding REPIN population do correlate 368 surprisingly well. RAYTs are absent from an entire S. maltophilia clade (middle of Figure 3A). This 369 clade has also lost a significant amount of green REPINs, and the proportion of the master 370 sequences in these populations is low (Figure 3C). Similarly, genomes without RAYTs have smaller 371 REPIN populations in the purple population than genomes with the corresponding RAYT (Figure 372 3D). A similar observation has been made previously in E. coli, P. chlororaphis and Neisseria 373 where the loss of the RAYT gene is followed by a decay of the REPIN population (Bertels, Rainey 374 2022).



sequences in these populations is low (Figure 4C). Similarly, genomes without orange RAYTs have
 smaller REPIN populations in the orange population than genomes with the corresponding RAYT
 (Figure 4D). A similar observation has been made previously in *E. coli, P. chlororaphis, N.* meningitidis and *N. gonorrhoeae* where the loss of the RAYT gene is followed by a decay of the

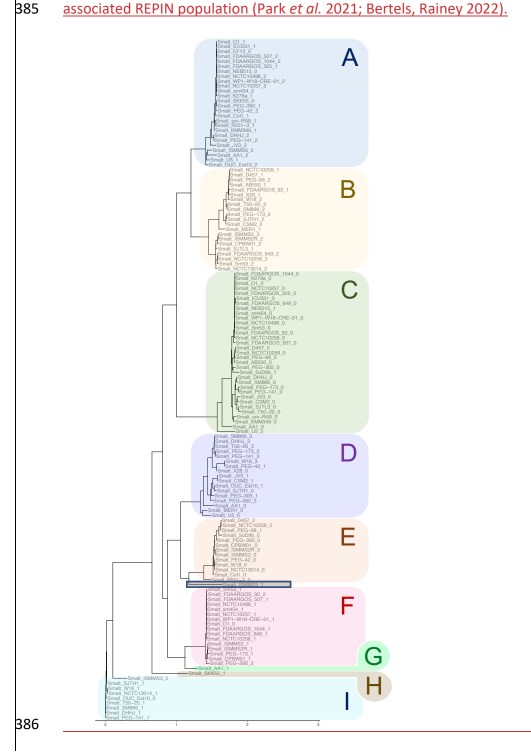


Figure 5. Phylogeny of RAYT genes and their associated REPINs. The tree shows RAYT genes from 49 *S. maltophilia* strains. Colours of clades A-I are assigned according to their association with a REPIN found within 130 bp **toof** the RAYT gene (see **Table 2**). Except for a single RAYT gene ISMMS3_1 (grey box), which could not be linked to a REPIN population.

RAYT population	REPIN palindromes
А	CCGACC AAC GGTCGG
В	CCAACC AA GGTTGGC
С	CCGGCC AGC GGCCGG
D	TCCACGC ATG GCGTGGA
E	CCGAGC CCAT GCTCGG
F	TCGACT AAC AGTCGA
G	TCGACC AAC GGTCGA
н	GCCGGGC ATG GCCCGGC
I	AGTCGAGC TT GCTCGACT

387 Table 2. REPIN palindromes associated with each RAYT clade.

Each RAYT clade from **Figure 45** is associated with a unique imperfect palindrome that is present at the 5' and/or 3' end of the REPIN (Figure 5). RAYT gene.

390

391 Linking REPIN populations with RAYT genes can be challenging

Unfortunately, RAREFAN is not always able to link the correct REPIN population with the correct RAYT gene. As shown in Figure 2In some RAREFAN runs indicate that sometimes associations between RAYTs and REPINs seem toare not be monophyletic, as for example the red RAYTs highlighted in red in Figure 2A3A. However, the same clade of RAYTs is uniformly coloured in yellow in Figure 2D3D, suggesting that the entire RAYT clade is associated with the same REPIN group. To investigate the true relationships between REPINs and RAYTs we first mapped REPIN groups to RAYT genes.

An analysis of all REPIN groups that were identified by RAREFAN across <u>fivefour</u> different RAREFAN runs <u>(Table 1, one additional analysis was performed with ISMMS3)</u> showed that there are a total of nine different REPIN groups, each defined by an individual central palindrome (**Table** 2). Each REPIN group is associated with a mono phyletic monophyletic RAYT group (Figure 4).
 There is only5). Only a single RAYT that we could is not identifyassociated with a REPIN for population (ISMMS3_1). Although there seems to be a one to one mapping between RAYT clades and REPIN groups, the question remains why RAREFAN sometimes links RAYT genes to the wrong REPIN group.

- RAREFAN could not link a REPIN to the RAYT gene ISMMS3 1 (Figure 5, grey box). While there is
 a sequence that resembles the A detailed analysispalindrome as well as variants of the extragenic
 spaceC palindrome flanking both sides of the RAYT gene (Supplementary Figure 2), none of the
 sequences formed REPIN populations large enough to be identified by RAREFAN. Presumably the
 RAYT ISMMS3_1, which is only present in a single *S. maltophilia* strain, is at the early stages of
 establishing a REPIN population "wrongly" associated, and the REPIN population has not spread
 to a considerable size yet.
- <u>There are two more cases where RAREFAN failed to link RAYT genes with any REPINs (ISSMS2</u>
 <u>and ISSMS2R 1, Supplementary Figure 1 D and E). Detailed sequence analyses showed that the</u>
 <u>respective REPINs are located at a distance of more than 130 bp (an adjustable parameter in</u>
 <u>RAREFAN). These REPINs are ignored by RAREFAN by default. However, this parameter can be</u>
 <u>adjusted manually and when set to a distance of 200 bp, RAREFAN correctly links these</u>
 <u>genesREPINs to the RAYT gene.</u>
 <u>In three cases the wrong REPIN population was linked to a RAYT gene. In our dataset this can</u>

421 happen when RAYTs are flanked by seed sequences from two different REPIN populations (Supplementary Figure 1 A-C). A single REP sequence from the "wrong" (non-monophyletic 422 423 RAYT) clade occurs together with multiple REP or REPIN sequences from the "right" 424 (monophyletic in a different RAREFAN run) clade. REPINs are linked to the "wrong" RAYT when the correct REPIN population is absent in the chosen reference genome. Moreover, the "wrongly" 425 426 associated REP singlets always show up as belonging to the REPIN population of a RAYT sister 427 group and show high sequence similarity This problem can be alleviated by performing analyses 428 with multiple reference genomes and comparing the results.

429 Additionally to linking the wrong REPINs and RAYTs, RAREFAN sometimes failed to link RAYT 430 genes with any REPINs. Detailed sequence analyses showed that in two out of a total of three 431 such cases the REPIN was located at a distance of more than 130bp (a parameter set in RAREFAN 432 that could be modified) (Supplementary Figure 1-D-E). REPINs that are located at a distance of 433 more than 130bp are ignored by RAREFAN. In a third case there was no REPIN that could be linked to the RAYT gene ISMMS3 1 (Figure 4, grey box). While there is a sequence that resembles the 434 A palindrome as well as variants of the C-palindrome flanking both sides of the RAYT gene, 435 436 (Supplementary Figure 2), none of the sequences formed REPIN populations large enough to be 437 identified by RAREFAN. Presumably the RAYT ISMMS3 1, which is only present in a single S. 438 maltophilia strain, is at the early stages of establishing a REPIN population. Based on our findings, 439 RAREFAN users should always critically analyse RAREFAN results, particularly when the results 440 require unusual evolutionary explanations.

441

Symmetric REPINs can be detected by RAREFAN:

Usually ~100 bp

Putative RAYT binding		Putative RAYT binding		
→ ←				
High sequence conservation	Low sequence conservation	High sequence conservation		
REP forward	REP reverse			

Asymmetric REPINs cannot be detected by RAREFAN:

_ → ←		→ ←	
Deletion		Insertion	

442

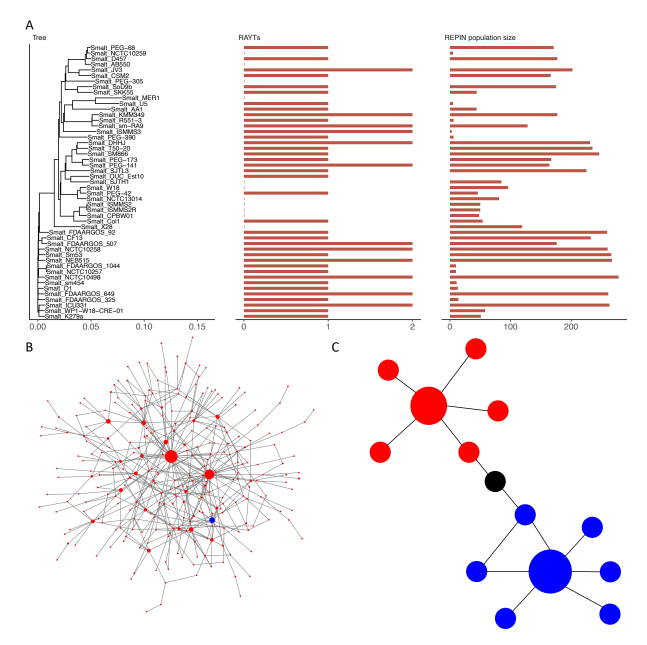
Figure 5. The structure of symmetric and asymmetric REPINs. A typical REPIN consists of two highly conserved regions at the 5' and 3' end of the REPIN (white), separated by a spacer region of lower sequence conservation (grey). The entire REPIN is a palindrome (blue arrows), which means it can form hairpin structures in single stranded DNA or RNA. Each 5' and 3' region contains a nested imperfect palindrome, which is referred to as REP (repetitive extragenic palindromic) sequence and has first been described in *E. coli* (Higgins *et al.* 1982). REPINs can be either symmetric or asymmetric. Asymmetric REPINs have a deletion and a corresponding insertion in the highly conserved 5' or 3' end, which leads to "bubbles" in the hairpin structure. REPINs in *E. coli* are asymmetric, which makes analyses with RAREFAN more challenging. Figure adapted from (Bertels, Rainey 2022).

444 REPIN groups may be lost when the seed distance is too large

The seed distance parameter determines whether two highly abundant sequences are sorted into the same or different REPIN groups (**Figure 182B**). If two REPINs from two different groups occur next to each other, at a distance of less than the seed distance parameter, then the two seeds are erroneously sorted into the same group. If two different REPIN groups are sorted into the same group then one of the groups will be ignored by RAREFAN, because only the most abundant seed <u>in each group</u> will be used to identify REPINs.

A manual analysis (*e.g.*, multiple sequence alignment) of sequences in the groupSeedSequences folder of the RAREFAN output can identify erroneously merged REPIN groups. In *S. maltophilia*, groups are separated well when the <u>distance</u> parameter is set to 15 <u>bp</u> and Sm53 is used as a reference. When the parameter is set to 30 <u>bp</u> instead, one of the REPIN groups will be missed by RAREFAN.

A small seed distance parameter will separate seed sequences belonging to the same REPIN
group into different groups. Hence, RAREFAN will analyse the same REPIN group multiple times.
While this will lead to increased RAREFAN runtimes, these errors, are easy to spot, because (1)
the same RAYT gene will be associated to multiple REPIN groups, (2) the central palindrome
between the group is identical and (3) the master sequence between the groups will be very
similar.



463

Figure 6. Closely related REPIN populations may be merged by RAREFAN. (A) REPIN group 2 identified in a *S. maltophilia* Sm53 RAREFAN run. The RAREFAN result suggests that REPIN group-2 is sometimes associated with two RAYTs. (**B**) A closer inspection of the data shows that Group 2 is a combination of two different REPIN groups, the <u>"real"</u> Group 2 and Group 0. The network shown, visualizes all REP sequences identified as Group 2. Nodes in the network represent 21 bp long REP sequences. Two nodes are connected if the sequences they represent differ by exactly one nucleotide. The node size indicates the abundance of the sequence in the genome. The blue node represents the most common Group 2 sequence, occurring 65 times in the genome. The largest red node occurs 407 times in the genome and resembles a Group 0 REP sequence (**C**) Illustration of how small changes to a single sequence can connect two sequence cluster. The most common 21 bp long sequence in Group 0 differs in only four positions from the

most common 21 bp long sequence in Group 2. There is a set of sequences that connects these two groups that only differ in exactly one position each (nodes connected by an edge), which passes through the black node. If there is such an unbroken path between REP sequences, then REPIN groups will be merged.

464

465 Closely related REPIN groups may be merged into a single group by RAREFAN

466 Incorrect merging of REPIN groups can occur when two REPIN groups are closely related. We 467 identified merged REPIN groups in S. maltophilia because RAREFAN linked some REPIN groups 468 with two RAYT genes in the same genome (Figure 6A). While REPIN groups linked to two RAYTs 469 has been observed before in Neisseria meningitidis (Bertels, Rainey 2022), it is particularly 470 unusual in S. While REPIN groups linked to two RAYTs has been observed before in Neisseria 471 meningitidis (Bertels, Rainey 2022), it is particularly unusual in S. maltophilia due to some key 472 differences between REPIN-RAYT in the two bacterial species. First, RAYTs in N. meningitidis 473 belong to Group 2 and RAYTs in S. maltophilia belong to Group 3 (Bertels, Gallie, et al. 2017), two 474 very divergent RAYT groups. Second, RAYTs that are associated to the same REPIN group in N. 475 meningitidis are almost identical, since they are copied by an insertion sequence in trans (Bertels, 476 Rainey 2022)meningitidis are almost identical, since they are copied by an insertion sequence in 477 trans (Bertels, Rainey 2022), something that is not the case for S. maltophilia, where the two 478 RAYTs are very distinct from each other (green and red clade in Figure 2A3A, or clade A and C in 479 Figure 45).

480 A closer inspection of all sequences identified in REPIN group 2 shows that it also contains 481 sequences belonging to REPIN group 0 (palindromes linked to clade A and C in **Table 2**). The relationship between the sequences shows that there is a chain of sequences that all differ by at 482 483 most a single nucleotide between the most abundant sequence in group 2 to the most abundant sequence in group 0 (Figure 6B and C). <u>Hence, the reason group 0 and group 2 are merged is that</u> 484 they are too closely related to each other and hybrids of the two REPIN groups exist. Because 485 486 sequence groups are built by identifying all related sequences in the genome recursively, closely 487 related groups (the REPIN group 0 seed only differs in four nucleotides from the REPIN group 2 488 seed sequence) can be merged into a single REPIN group. REPIN population size and RAYT number 489 are the sum of REPIN group 0 and 2. There are various possibilities to resolve this issue: (1)

490 subtract sequences from group 0 (which does not contain group 2) from REPIN group 2; (2) use 491 a different sequence seed from the group 2 seed collection in the seed sequence file 492 (groupSeedSequences/Group_Smalt_Sm53_2.out); or (3) sometimes it may be possible 493 to rerun RAREFAN with a different reference strain where the issue does not occur; or (4) increase 494 the length of the seed sequence.

495

496 <u>Performance</u>

We measured the elapsed time for a complete RAREFAN run for three different species and for
 5, 10, 20, and 40 genomes with randomly selected reference strains and the two query RAYTs
 (yafM Ecoli and yafM SBW25). For a given number N of submitted genomes of average
 sequence length L (in megabases), a RAREFAN run completes in approximately T = (8-10 seconds)
 * N * L on our moderate server hardware (4 CPU cores, 16 GB shared RAM) (Supplementary
 Figure 3 and 4).

503 Discussion

RAREFAN allows users to quickly detect REPIN populations and RAYT transposases inside
bacterial genomes. It also links the RAYT transposase genes to the REPIN population it duplicates.
These data help the user to study REPIN-RAYT dynamics in their strains of interest without a
dedicated bioinformatician, and hence will render REPIN-RAYT systems widely accessible.

508 One limitation of RAREFAN is that REPINs can only be identified in genomes when they are 509 symmetric (Figure 51). Symmetric REPINs have seed sequences that can morph into each other 510 by a series of single substitutions (intermediate sequences need to be present in the genome). A 511 REPIN consists of a 5' and a 3' REP sequence. If one of these REP sequences contains an insertion 512 or deletion, which the other REP sequence does not contain then RAREFAN will not recognize the 513 second repeat of the seed sequence. In this case, RAREFAN will not be able to identify REPINs but 514 can still be used to analyze REP singlet populations. To date, the only known asymmetric REPIN 515 population populations are found in *E. coli* REPINS. However, it is likely that asymmetric REPINs 516 also exist in other microbial species.

RAREFAN sometimes cannot correctly divide REPINs into REPIN groups. Either because REPINs
from different groups occur in close proximity in the genome, an issue that can easily be solved
by adjusting a RAREFAN parameter, or because two REPIN groups are very closely related (Figure **6**). Unfortunately, RAREFAN is not able to automatically detect and resolve the assignment of
closely related REPINs into groups yet. Hence it is advisable to manually check associations
between REPIN groups and RAYT genes by analyzing the composition of REPIN groups.

523 In the future we aim to make RAREFAN even more versatile and easier to use by, for example, 524 automatically integrating data from public databases such as Genbank, and integrating RAREFAN 525 into workflows such as Galaxy (Afgan *et al.* 2018).

526 RAREFAN makes the study of REPIN-RAYT systems more accessible to any biologist or 527 bioinformatician interested in studying intragenomic sequence populations. Our tool will help 528 understand the purpose and evolution of REPIN-RAYT systems in bacterial genomes.

529 Acknowledgements

We would like to thank Prajwal Bharadwaj for assisting us with the sequence analysis and JennaGallie for valuable feedback on the manuscript.

532 References

533 Afgan E, Baker D, Batut B, van den Beek M, Bouvier D, Cech M, Chilton J, Clements D, Coraor N,

534 Grüning BA, Guerler A, Hillman-Jackson J, Hiltemann S, Jalili V, Rasche H, Soranzo N,

535 Goecks J, Taylor J, Nekrutenko A, Blankenberg D (2018) The Galaxy platform for

536 accessible, reproducible and collaborative biomedical analyses: 2018 update. *Nucleic*

537 Acids Res., 46, W537-W544-W537–W544. https://doi.org/10.1093/nar/gky379

538 Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ (1990) Basic local alignment search tool.

539 *Journal of Molecular Biology*, **215**, 403–410. https://doi.org/10.1006/jmbi.1990.9999

540 Arnold K, Gosling J, Holmes D (2005) The Java programming language. Addison Wesley

541 Professional.

542 Bertels F, Gallie J, Rainey PB (2017) Identification and Characterization of Domesticated Bacterial

- 543 Transposases. *Genome Biology and Evolution*, **9**, 2110–2121. 544 https://doi.org/10.1093/gbe/evx146
- 545 Bertels F, Gokhale CS, Traulsen A (2017) Discovering Complete Quasispecies in Bacterial 546 Genomes. *Genetics*, **206**, 2149–2157. https://doi.org/10.1534/genetics.117.201160
- 547 Bertels F, Rainey PB (20112011a) Within-Genome Evolution of REPINs: a New Family of Miniature
- 548 Mobile DNA in Bacteria. *PLoS genetics*, **7**, e1002132. 549 https://doi.org/10.1371/journal.pgen.1002132
- Bertels F, Rainey PB (2011b) Curiosities of REPINs and RAYTs. *Mobile Genetic Elements*, 1, 262–
 268. https://doi.org/10.4161/mge.18610
- 552 Bertels F, Rainey PB (2022) Ancient Darwinian replicators nested within eubacterial genomes. ,
- 553 2021.07.10.451892. https://doi.org/10.1101/2021.07.10.451892
- 554 Bichsel M, Barbour AD, Wagner A (2010) The early phase of a bacterial insertion sequence 555 infection. *Theoretical Population Biology*. https://doi.org/10.1016/j.tpb.2010.08.003
- 556 Camacho C, Coulouris G, Avagyan V, Ma N, Papadopoulos J, Bealer K, Madden TL (2009) BLAST+:
- 557 architecture and applications. *BMC Bioinformatics*, **10**, 421–9.
- 558 https://doi.org/10.1186/1471-2105-10-421
- van Dijk B, Bertels F, Stolk L, Takeuchi N, Rainey PB (2022) Transposable elements promote the
- 560 evolution of genome streamlining. *Philosophical Transactions of the Royal Society B:*
- 561 *Biological Sciences*, **377**, 20200477. https://doi.org/10.1098/rstb.2020.0477
- 562 Edgar RC (2004) MUSCLE: multiple sequence alignment with high accuracy and high throughput.
- 563 *Nucleic Acids Research*, **32**, 1792–1797. https://doi.org/10.1093/nar/gkh340

- 564 <u>Felsenstein J (1985) Phylogenies and the comparative method. *American Naturalist*, 1–15.</u>
- 565 Grinberg M (2018) *Flask web development: developing web applications with python*. O'Reilly 566 Media, Inc.
- 567 Guindon S, Dufayard J-F, Lefort V, Anisimova M, Hordijk W, Gascuel O (2010) New algorithms and
- 568 <u>methods to estimate maximum-likelihood phylogenies: assessing the performance of</u>
- 569 PhyML 3.0. Systematic Biology, **59**, 307–321. https://doi.org/10.1093/sysbio/syq010
- 570 Haubold B, Klötzl F, Pfaffelhuber P (2015) andi: fast and accurate estimation of evolutionary
- 571 distances between closely related genomes. *Bioinformatics*, **31**, 1169–1175.
- 572 https://doi.org/10.1093/bioinformatics/btu815
- 573 Higgins CF, Ames GF, Barnes WM, Clement JM, Hofnung M (1982) A novel intercistronic
- 574 regulatory element of prokaryotic operons. *Nature*, **298**, 760–762.
 575 https://doi.org/10.1038/298760a0
- 576 Initiative TOS (2021) The MIT License.
- 577 Kearse M, Moir R, Wilson A, Stones-Havas S (2012) Geneious Basic: an integrated and extendable
- 578 desktop software platform for the organization and analysis of sequence data.
- 579 Kleinmann SG, Rudolph S, Vila S, Rodin J, Peña JF-S (2021) *The Debian GNU/Linux Operating* 580 *System Manual*.
- Lawrence JG, Ochman H, Hartl DL (1992) The evolution of insertion sequences within enteric
 bacteria. *Genetics*, **131**, 9–20. https://doi.org/10.1093/genetics/131.1.9
- 583 Nunvar J, Huckova T, Licha I (2010) Identification and characterization of repetitive extragenic
 584 palindromes (REP)-associated tyrosine transposases: implications for REP evolution and

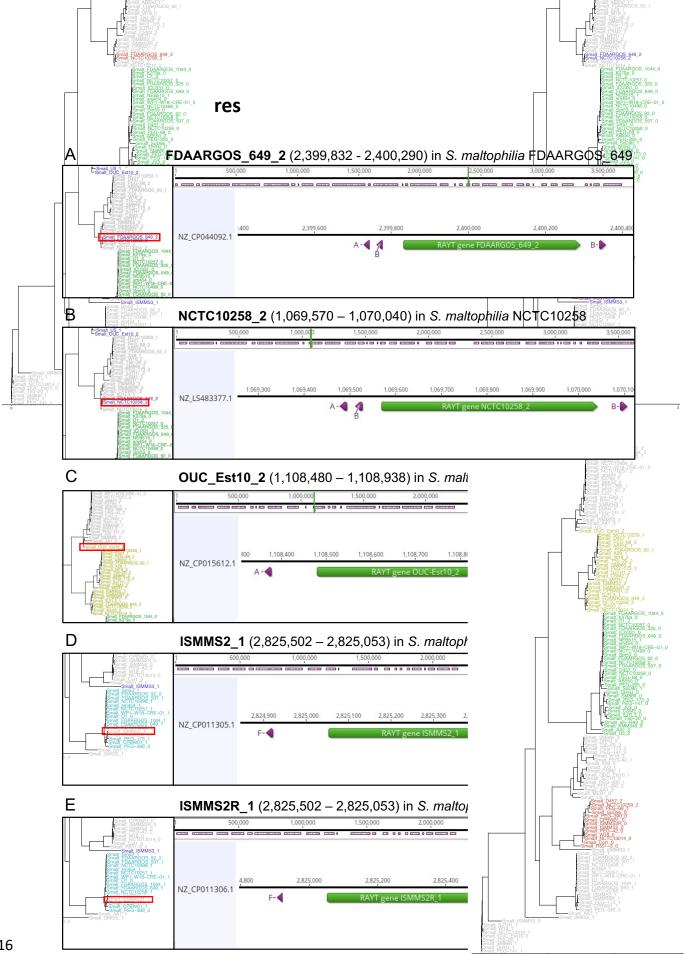
- 585 dynamics in bacterial genomes. *BMC Genomics*, **11**, 44. https://doi.org/10.1186/1471-586 2164-11-44
- 587 Park HJ, Gokhale CS, Bertels F (2021) How sequence populations persist inside bacterial genomes.
 588 *Genetics*, **217**. https://doi.org/10.1093/genetics/iyab027
- 589 R Core Team (2016) R: A Language and Environment for Statistical Computing. R Foundation for
 590 Statistical Computing, Vienna, Austria.
- 591 Rankin DJ, Bichsel M, Wagner A (2010) Mobile DNA can drive lineage extinction in prokaryotic
- 592 populations. Journal of Evolutionary Biology. https://doi.org/10.1111/j.1420-
- 593 9101.2010.02106.x
- 594 RStudio, Inc (2013) Easy web applications in R.
- 595 Rutherford K, Parkhill J, Crook J, Horsnell T, Rice P, Rajandream MA, Barrell B (2000) Artemis:
- 596 sequence visualization and annotation. *Bioinformatics*, 16, 944–945.
 597 https://doi.org/10.1093/bioinformatics/16.10.944
- 598 Sawyer SA, Dykhuizen DE, DuBose RF, Green L, Mutangadura-Mhlanga T, Wolczyk DF, Hartl DL
- 599 (1987) Distribution and Abundance of Insertion Sequences Among Natural Isolates of
- 600 Escherichia coli. *Genetics*, **115**, 51–63. https://doi.org/10.1093/genetics/115.1.51
- Ton-Hoang B, Siguier P, Quentin Y, Onillon S, Marty B, Fichant G, Chandler M (2012) Structuring
- the bacterial genome: Y1-transposases associated with REP-BIME sequences. *Nucleic Acids Research*, 40, 3596–3609. https://doi.org/10.1093/nar/gkr1198
- 604 Van Dongen S (2000) A cluster algorithm for graphs. *Report-Information systems*, 1–40.
- 605 Van Rossum G, Drake Jr FL (1995) *Python reference manual*. Centrum voor Wiskunde en 606 Informatica Amsterdam.

transposition bursts help the elements persist? *BMC Evolutionary Biology*, 15, 288–12.
https://doi.org/10.1186/s12862-015-0560-5
Yu G, Lam TT-Y, Zhu H, Guan Y (2018) Two Methods for Mapping and Visualizing Associated Data
on Phylogeny Using Ggtree. (FU Battistuzzi, Ed,). *Molecular biology and evolution*, 35,
3041–3043. https://doi.org/10.1093/molbev/msy194

Wu Y, Aandahl RZ, Tanaka MM (2015) Dynamics of bacterial insertion sequences: can

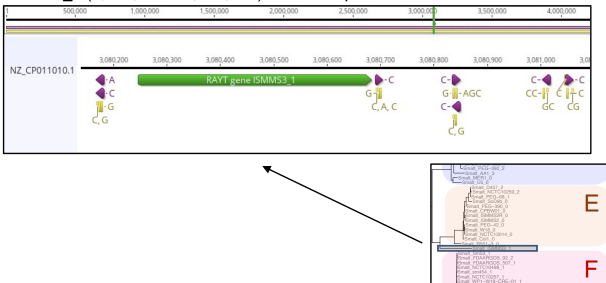
613

607



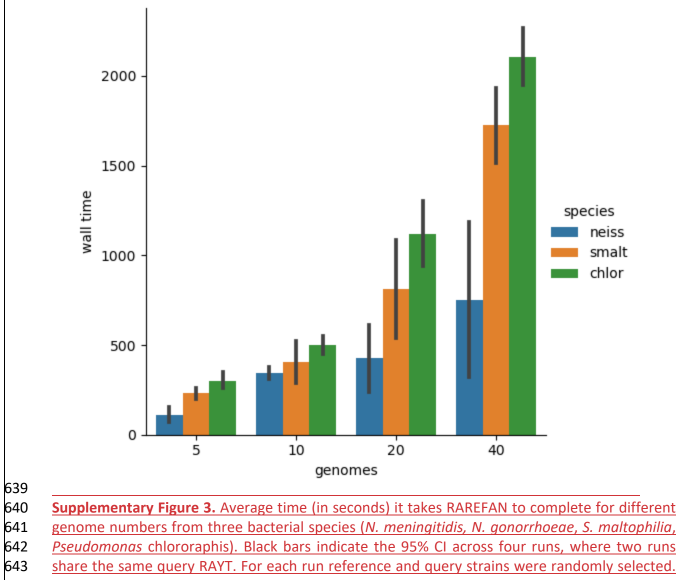
617 Supplementary Figure 1:. Sequence analysis shows REPIN groups are indeed associated with 618 monophyletic RAYTs. Non-monophyletic or missing associations to REPIN populations identified 619 by RAREFAN were investigated in the corresponding genomes using Geneious (Kearse et al. 620 2012). Red boxes mark the position of the atypical RAYT that is being analyzed in detail. Mapping 621 of REPIN palindromes A-I (with zero mismatches) shows FDAARGOS 649 2 (A), NCTC10258 2 622 (B), and OUC Est 2 (C) are linked to the wrong REPIN group because REP singlets that are 623 ordinarily linked to a RAYT sister clade are found in close proximity to the RAYT. These wrong 624 associations between REPIN and RAYT usually occur when the correct REPIN population is absent 625 from the reference genome. ISMMS2R 1 (D) and ISMMS2 1 (E) were not linked to REPIN 626 population by RAREFAN because the corresponding seed sequences were located at a distance 627 of more than 130 bp from the RAYT gene. -Nucleotide sequences and positions were extracted from output files generated by RAREFAN. Complete genome sequences are available in NCBI 628 629 Nucleotide Database using Accessions: (A) NZ CP044092.1, (B) NZ LS483377.1, (C) 630 NZ CP015612.1, (D) NZ CP011306.1, (E) NZ CP011305.1.

ISMMS3_1 (3,080,683 - 3,080,246) in S. maltophilia ISMMS3



631

Supplementary Figure 2: RAYT gene ISMMS3_1 cannot be linked to a REPIN population. The sequence of the RAYT gene ISMMS3_1 and its flanking sequences was analyzedwere analysed in Geneious (Kearse *et al.* 2012). The inset shows the location of ISMMS3_1 in the RAYT phylogeny (grey box). When mapping all of the identified palindromes to the RAYT region and allowing up to four mismatches (yellow annotations), various mutants of palindrome C were found in close proximity of the RAYT gene. However, we could not identify a corresponding REPIN population, which may indicate that the population has not yet expanded in the genome.



- 644 <u>All measurements were performed on 4CPU cores with 16 GB of shared memory.</u>
- 645

