# 1 Analysis of next- and third-generation RNA-Seq data reveals the structures of

# 2 alternative transcription units in bacterial genomes

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# 12 ABSTRACT

13 Alternative transcription units (ATUs) are dynamically encoded under different conditions or

14 environmental stimuli in bacterial genomes, and genome-scale identification of ATUs is essential for

15 studying the emergence of human diseases caused by bacterial organisms. However, it is unrealistic to

- 16 identify all ATUs using experimental techniques, due to the complexity and dynamic nature of ATUs.
- 17 Here we present the first-of-its-kind computational framework, named SeqATU, for genome-scale ATU
- 18 prediction based on next-generation RNA-Seq data. The framework utilizes a convex quadratic

19	programming model to seek an optimum expression combination of all of the to-be-identified ATUs.
20	The predicted ATUs in <i>E. coli</i> reached a precision of 0.77/0.74 and a recall of 0.75/0.76 in the two RNA-
21	Sequencing datasets compared with the benchmarked ATUs from third-generation RNA-Seq data. We
22	believe that the ATUs identified by SeqATU can provide fundamental knowledge to guide the
23	reconstruction of transcriptional regulatory networks in bacterial genomes.
24	INTRODUCTION
25	An operon in bacterial genomes is defined as a group of consecutive genes regulated by a common
26	promoter that all share the same terminator $(1)$ . Genes in the same operon generally encode proteins
27	with relevant or similar biological functions; e.g., <i>lacZ</i> , <i>lacY</i> , and <i>lacA</i> in the <i>lac</i> operon encode proteins
28	that help cells use lactose $(1, 2)$ . With decades of research on bacterial transcriptional regulation, the
29	operon model has been found to have complex mechanisms that control expression $(3-5)$ . Multiple
30	studies have shown that bacterial genes are dynamically transcribed under different triggering
31	conditions, leading to shared genes among different mRNA transcripts (6-8). This dynamic architecture
32	can be redefined by all of the alternative transcription units (a.k.a., ATUs) $(3, 5)$ , and more details can be
33	found in fig. S1.
34	ATU identification is of fundamental importance for understanding the transcriptional regulatory
35	mechanisms of bacteria, and these dynamic structures have been demonstrated to be associated with
36	human diseases (9-12). For example, Bhat et al. studied the alr-groEL1 operon, which is essential for the

37 survival or virulence of *M. tuberculosis* (9, 11), the causative agent of tuberculosis (TB), and found that

38	the regulation of the sub-operon is distinct from the main operon ( <i>alr-groEL1</i> operon) under stress,
39	especially during heat shock, pH, and SDS stresses (9). Another example is Helicobacter pylori, a
40	gastric pathogen that is the primary known risk factor for gastric cancer (12). Sharma et al. found an
41	acid-induced sub-operon cag22-18 transcribed from the primary cag25-18 operon in the cag
42	pathogenicity island of the <i>H. pylori</i> genome under acid stress (10). The mechanism of the complex ATU
43	structure in these pathogenic bacteria can help us to study the emergence of human diseases caused by
44	bacterial organisms.
45	Several newly developed techniques have provided a comprehensive view of the E. coli
46	transcriptome by identifying full-length primary transcripts (13-17). For example, SMRT-Cappable-seq
47	(6) combines the isolation of the full-length bacterial primary transcriptome with PacBio SMRT (Single
48	Molecule, Real-Time) sequencing (6), and simultaneous 5' and 3' end sequencing (SEnd-seq) (7)
49	captures both transcription start sites (TSSs) and transcription termination sites (TTSs) via
50	circularization of transcripts (17). Despite the great progress in experimental techniques, there are still
51	some deficiencies. On the one hand, the read depth and error rate of the third-generation sequencing
52	used in SMRT-Cappable-seq have an impact on ATU prediction compared with Illumina-based RNA-
53	Seq (7, 18). On the other hand, the time-consuming, laborious, and costly properties of these
54	experimental techniques make them unrealistic to be generally applicable to ATU predictions in bacteria
55	under specific conditions. Thus, novel and robust computational methods for ATU identification in
56	bacterial genomes based on RNA-Seq are urgently needed.

57	Fortunately, many computational studies have been carried out to predict ATUs in bacteria, which
58	have provided some preliminary studies for ATU prediction. Several public databases, such as
59	RegulonDB (19), DBTBS(20), MicrobesOnline (21), DOOR (22, 23), OperomeDB (24), DMINDA 2.0
60	(25), and ProOpDB (26), provide various levels of operon information and small amounts of ATU
61	information. However, these databases cannot provide genome-scale ATU information under specific
62	conditions. Some computational studies, including Rockhopper (27), SeqTU (4, 28), BAC-
63	BROWSER(29), rSeqTU (5), and Operon-mapper (30), utilize machine learning and model integration
64	methods based on genomic information and gene expression profiles to identify bacterial transcription
65	architecture. However, these works still cannot solve the dynamic patterns and overlapping nature of
66	ATUs.
67	Here, we present SeqATU, a novel computational method for genome-scale ATU prediction by
68	analyzing next- and third-generation RNA-Seq data (Fig. 1 and table S1). SeqATU utilizes a convex
69	quadratic programming model (CQP) and aims to provide the optimum expression combination of all of
70	the to-be-identified ATUs. Specifically, CQP minimizes the squared error between the predicted
71	expression level of ATUs and the actual expression levels in genetic and intergenic regions. It is
72	noteworthy that SeqATU also utilizes the information about the bias rate function in modeling non-
73	uniform read distribution as the linear constraints of CQP to profile the complexity of the ATU
74	architecture. Overall, SeqATU provides a generalized framework for the inference of ATUs based on
75	next-generation RNA-Seq data collected under multiple conditions and can be easily applied to any

#### <sup>76</sup> bacterial organism to identify the ATU architecture and construct a transcriptional regulatory network.

77

Please place Fig. 1 here.

#### 78 MATERIALS AND METHODS

# 79 Data collection

80 The two Cappable RNA-Seq datasets used in this study, M9Enrich Seq and RiEnrich Seq, were obtained from E. coli grown under two different conditions: M9 minimal medium and Rich medium, 81 respectively (6). The full-length primary transcripts were enriched as described in (6) with modifications 82 83 to be adapted to Illumina sequencing. The capping and polyA tailing were performed as described in (6). The capped RNA was enriched using hydrophilic streptavidin magnetic beads (New England Biolabs) 84 and eluted with Biotin using the same condition (6). Differently, the eluted RNA was enriched once 85 more using streptavidin beads to further remove processed RNA (e.g., rRNA). Subsequently, the eluted 86 87 RNA was used for library preparation using NEBNext Ultra II directional RNA library prep kit (E7760). Sequencing was performed on the Illumina Miseq system (paired-end, 100bp). All reads were mapped to 88 the *E. coli* genome using Burrows-Wheeler Aligner (BWA) with the default parameters (31). Read 89 90 alignment and other computational analyses were carried out using the E. coli genome NC 000913.3, 91 and the corresponding gene annotations (GCF 000005845.2 ASM584v2 genomic.gff) were downloaded from NCBI. Two experimentally verified ATU datasets, SMRT M9Enrich and 92 93 SMRT RiEnrich, were used as the benchmark data to evaluate the predicted ATUs, which were

generated by SMRT-Cappable-seq under the same conditions as the Illumina datasets M9Enrich\_Seq
and RiEnrich\_Seq, respectively (6). In addition, the ATUs defined by RegulonDB (19) and SEnd-seq (7)
were also used as additional evaluation data in our study.

#### 97 Calculation of the expression values of genetic and intergenic regions

After the RNA-Seq reads in M9Enrich\_Seq and RiEnrich\_Seq were mapped to the *E. coli* genome using BWA, we determined the number of reads N(l) covering each genomic position *l*. Suppose that  $g_i$ and  $g_{i+1}$  are two consecutive genes on the same strand; we denote the expression value of  $g_i$  as  $c_i$ and the expression value of the intergenic region between genes  $g_i$  and  $g_{i+1}$  as  $b_{i,i+1}$ . Then, the calculation of  $c_i$  and  $b_{i,i+1}$  is given by:

$$c_i = \frac{\sum_{k \in g_i} N(k)}{|g_i|} \tag{1}$$

$$b_{i,i+1} = \frac{\sum_{l \in g_{i,i+1}} N(l)}{|g_{i,i+1}|} \tag{2}$$

103 where  $k \in g_i$  denotes that genomic position k is on the gene  $g_i$  and  $|g_i|$  denotes the genomic 104 length of  $g_i$ .

# 105 Modeling non-uniform read distribution along mRNA transcripts

We introduced the bias rate function, which is similar to the bias curves in the work of Wu *et al.* (*32*), to address the non-uniform distribution of the RNA-Seq reads along mRNA transcripts (*32-35*). The bias function reflects the relative read distribution bias from the 3' end to the 5' end of an mRNA transcript.

109	We assumed that the maximum read coverage of all the genomic positions of an mRNA transcript is the
110	expression level without bias. It is noteworthy that a single gene mRNA transcript with no shared gene
111	among different mRNA transcripts can serve as the ideal template for modeling non-uniform read
112	distribution along mRNA transcripts. The specific steps of modeling non-uniform read distribution are
113	detailed as follows:

Step 1: Single Gene mRNA Transcript Selection. We selected single gene mRNA transcripts from the 114 evaluation data and plotted their expression distributions. Specifically, 12 groups of single gene mRNA 115 transcripts with lengths ranging from 300 to 1,500 bp were selected from the evaluation data (more 116 details are given in method S1), and each group had ten randomly chosen mRNA transcripts. Apparent 117 decline trends appeared in the single gene mRNA transcripts with long lengths, ranging from 1,100 to 118 1,500 bp (fig. S2). The reason for this phenomenon may be that the incomplete transcription and 3' end 119 120 degradation or processing induce the enrichment of signal at 5' end of the mRNA transcripts with long lengths (36, 37). Finally, we plotted the expression distribution of single gene mRNA transcripts with 121 lengths ranging from 1,100 to 1,500 bp. 122

123 Step 2: Acquiring the Bias Rate Function. We applied nonlinear regression to the expression

distribution of the selected single gene mRNA transcripts and acquired the hypothetical function f(x). Specifically, the x axis and y axis of the expression distribution were converted to the distance from the 3' end of an mRNA transcript and the bias rate of read distribution, respectively. To apply nonlinear regression to single gene mRNA transcripts with different lengths, normalization was also implemented

128 on x. Here,  $x = (x_1, x_2, ..., x_m)$  and  $y = (y_1, y_2, ..., y_m)$  are defined by:

$$x_{i} = \begin{cases} \frac{l_{m} - l_{m-i+1}}{max_{l} - l_{1}} \times 10^{3}, & forward \\ \frac{l_{i} - l_{1}}{max_{l} - l_{1}} \times 10^{3}, & reverse \end{cases}$$
(3)

$$y_{i} = \begin{cases} \frac{N(l_{m-i+1})}{max_{y}}, & forward\\ \frac{N(l_{i})}{max_{y}}, & reverse \end{cases}$$
(4)

129 where *m* denotes the number of genomic positions on an mRNA transcript;  $l = (l_1, l_2, ..., l_m)$  denotes 130 the genomic positions on an mRNA transcript;  $max_l = l_m$ ;  $N(l_i)$  denotes the expression level of the 131 genomic position  $l_i$ , i.e., the number of reads covering the genomic position  $l_i$ ; and  $max_y$  denotes the 132 expression level without bias in an mRNA transcript, which is calculated as  $max \{N(l_i)\}, 1 \le i \le m$ . 133 We used the function *nls* in R to acquire the hypothetical function f(x).

Step 3: Constructing Bias Rate Vectors. We constructed a genetic or intergenic region bias rate vector
 for each mRNA transcript by calculating the bias rate of all of its component genetic or intergenic

136 regions. The bias rate of a genetic or an intergenic region is the average bias rate of all the genomic

137 positions that it contains. Considering an mRNA transcript T and its component gene set

138  $\{g_1, g_2, \dots, g_n\}$  (the details of the gene labels are described in method S2), we denoted the genetic

139 region bias rate vector as  $\mathbf{u} = (u_1, u_2, ..., u_n)$ , which was calculated using the formula:

$$u_{i} = \begin{cases} \frac{\sum_{t=m-i_{q}+1}^{m-i_{p}+1} f(x_{t})}{x_{m-i_{p}+1} - x_{m-i_{q}+1} + 1}, & forward\\ \frac{\sum_{t=i_{p}}^{i_{q}} f(x_{t})}{x_{i_{q}} - x_{i_{p}} + 1}, & reverse \end{cases}$$
(5)

where *m* denotes the number of genomic positions on *T*;  $u_i$  denotes the bias rate of  $g_i$  for *T*; and  $L_g = (l_{1p}, l_{1q}, l_{2p}, l_{2q}, ..., l_{np}, l_{nq})$  is the range of the genomic positions of  $\{g_1, g_2, ..., g_n\}$ , while the range of the genomic positions of  $g_i$  is  $[l_{ip}, l_{iq}]$ ,  $1 \le i \le n$ . Similarly, the calculation of the intergenic region bias rate vector  $\boldsymbol{v} = (v_1, v_2, ..., v_{n-1})$  is provided in method S3.

## 144 Modification of maximal ATU clusters

A maximal ATU cluster is defined as a maximal consecutive gene set such that each pair of its 145 consecutive genes can be covered by at least one ATU. Similar to ATUs, maximal ATU clusters are also 146 dynamically composed under different conditions or environmental stimuli in bacterial genomes (5, 38). 147 Such a maximal ATU cluster can be used as an independent genomic region for ATU prediction, which 148 alleviates the difficulty in computationally predicting ATUs at the genome scale. The output of our in-149 house tool rSeqTU can serve as the maximal ATU cluster data, which lays a solid foundation for ATU 150 prediction (5). We modified the maximal ATU clusters from rSeqTU: (i) two maximal ATU clusters with 151 152 distances less than 40 bp were combined into one cluster and (ii) a maximal ATU cluster was split at the 153 intergenic region where the opposite-strand genes were located. In addition, we selected the maximal ATU clusters with expression values over ten (see the details in method S4), according to the study of 154 Etwiller et al. (13). 155

# 156 The mathematical programming model for ATU prediction

157 The predicted ATU expression profile should be consistent with the observed expression profiles of the

genetic and intergenic regions. Therefore, the prediction of the ATU profiles can be modeled as an
optimization problem, which seeks an optimum expression combination of all of the to-be-identified
ATUs to minimize the gap between the predicted ATUs and the observed genetic and intergenic region
expression profiles. Here, a convex quadratic programming model was built to solve this optimization
problem.

163 We denoted a maximal ATU cluster as G, assuming that it contains the consecutive genes

164  $\{g_1, ..., g_n\}$ , and the intergenic regions of these genes are  $\{g_{1,2}, ..., g_{n-1,n}\}$ . The size of *G* is defined as 165 the number of its component genes *n*. Theoretically, there are  $\frac{n \times (n+1)}{2}$  ATUs for *G*, and an ATU with 166 consecutive genes  $\{g_i, g_{i+1}, ..., g_j\}$  is denoted as  $a^{i,j}$ ; the corresponding expression value is  $x^{i,j}$ ,  $1 \le$ 167  $i \le j \le n$ .

For the component gene  $g_k$  of G, the gap between the gene expression value  $c_k$  and the sum of the 168 expression level of the ATUs containing it is denoted as  $\tau_k$ , which provides the first n equality 169 constraints in our mathematical programming model, k = 1, 2, ..., n. Similarly, for the intergenic region 170  $g_{l,l+1}$  of G, the gap between the intergenic region expression value  $b_{l,l+1}$  and the sum of the 171 expression level of the ATUs containing it is denoted as  $\beta_l$ , providing the last n-1 equality 172 173 constraints in our mathematical programming model, l = 1, 2, ..., n - 1. The goal of our mathematical programming model is to minimize the square of  $\varepsilon =$ 174  $(\tau_1, \tau_2, ..., \tau_n, \beta_1, ..., \beta_{n-1})$ , as the combination of  $x^{i,j}$  with a minimal value of  $\epsilon \epsilon^T$  is corresponding to 175 an optimum expression combination of all ATUs  $a^{i,j}$  for  $G, 1 \le i \le j \le n$ . Additionally, to control the 176

177 number of optimal solutions and reduce the false-positive errors, we added an  $L^1$  regularization  $\alpha ||\mathbf{x}||_1$ 178 to  $\boldsymbol{\varepsilon}\boldsymbol{\varepsilon}^T$  with  $x^{i,j} \ge 0$ , which is a linear function. Because of the variant expression level of different 179 maximal ATU clusters, we used the expression value of G as  $\alpha$ . In total, the convex quadratic 180 programming model with unknown variables  $(\mathbf{x}, \boldsymbol{\varepsilon})$  is shown as follows:

$$min \qquad \varepsilon \varepsilon^{T} + \alpha ||\mathbf{x}||_{1}$$
s.t. 
$$\sum_{i=1}^{k} \sum_{j=k}^{n} u_{i,k} x^{i,j} = c_{k} + \tau_{k} \qquad k = 1, 2, ..., n$$

$$\sum_{i=1}^{l} \sum_{j=l+1}^{n} v_{i,l+1} x^{i,j} = b_{l,l+1} + \beta_{l} \qquad l = 1, 2, ..., n - 1$$

$$\mathbf{x} = (x^{i,j}), \qquad x^{i,j} \ge 0 \qquad 1 \le i \le j \le n$$

$$\varepsilon = (\tau_{1}, \tau_{2}, ..., \tau_{n}, \beta_{1}, ..., \beta_{n-1}) \qquad (6)$$

181 where  $\boldsymbol{u} = (u_{i,j})$  is the genetic region bias rate vector for G,  $u_{i,j}$  is the bias rate of gene  $g_j$  for ATU 182  $a^{i,k}$ ,  $1 \le i \le j \le n$ ,  $j \le k \le n$ ,  $\boldsymbol{v} = (v_{p,q})$  is the intergenic region bias rate vector for G, and  $v_{p,q}$ 183 is the bias rate of the intergenic region  $g_{q-1,q}$  for ATU  $a^{p,l}$ ,  $1 \le p < q \le n$ ,  $q \le l \le n$  (see the 184 details in method S5).

# 185 **Two evaluation methods for ATU prediction**

In the first evaluation method, precision and recall were defined based on perfect matching (Eqs. 7). Perfect matching of two ATUs means that all of their component genes are the same. Here, the true positives (*TP*) are the number of predicted ATUs with the same component genes as an ATU in the evaluation data; the false positives (*FP*) are the number of predicted ATUs that do not exist in the

evaluation data; the false negatives (FN) are the number of ATUs that appear in the evaluation data but

191 not in the prediction results of SeqATU.

$$precision = \frac{TP}{TP + FP}$$
$$recall = \frac{TP}{TP + FN}$$
(7)

In the second evaluation method, precision and recall were defined based on relaxed matching, which 192 is measured by the similarity of two ATUs. Assuming that an ATU t is in one of two datasets (the 193 predicted ATU dataset and evaluated ATU dataset), the definition and calculation of the similarity of t 194 are shown in the following three cases: 195 Case 1: If t shares boundary genes at both ends of an ATU in the other dataset, i.e., all component 196 genes of t are the same as one in the other dataset, then similarity(t) = 1. 197 Case 2: If t shares exactly one boundary gene of ATUs in the other dataset, then we denote  $U_a$  as 198 the ATUs in the other dataset that share the 5'-end gene with t and denoted  $U_b$  as the ATUs in the 199 other dataset that share the 3'-end gene with t,  $U_a \cap U_b = \emptyset$ , one of  $U_a$  and  $U_b$  can be empty. Then, 200 similarity(t) =  $\frac{1}{2} \max_{t' \in U_a} \frac{\alpha(t')}{\beta(t')} + \frac{1}{2} \max_{t' \in U_b} \frac{\alpha(t')}{\beta(t')}$ (8) where  $\alpha(t')$  is the number of shared genes of t and t' and  $\beta(t')$  is the maximal size of t and t'. 201 Case 3: If t shares no boundary genes at both ends of the ATUs in the other dataset, then 202

203 similarity(t) = 0.

Finally, the precision and recall based on relaxed matching are calculated by the following formula:

$$precision = \frac{\sum_{t \in T_1} similarity(t)}{n_1}$$

$$recall = \frac{\sum_{t \in T_2} similarity(t)}{n_2} \tag{9}$$

where  $T_1$  is the set of predicted ATUs,  $n_1$  is the number of predicted ATUs,  $T_2$  is the set of evaluated ATUs, and  $n_2$  is the number of evaluated ATUs.

207 **RESULTS** 

# A reliable bias rate function is acquired in modeling non-uniform read distribution along mRNA transcripts

210 To ensure the reliability of the bias rate function in modeling non-uniform read distribution, we selected

211 four single gene mRNA transcript datasets randomly from the two evaluation datasets

212 (SMRT\_M9Enrich and SMRT\_RiEnrich), named M9Enrich\_1, M9Enrich\_2, RiEnrich\_1, and

213 RiEnrich\_2. Four bias rate functions, which are exponential functions, were generated after conducting

214 nonlinear regression on the mRNA transcripts across these four datasets (Fig. 2). We found that these

bias rate functions were similar ( $R^2 > 0.998$ ) when we evaluated the R-square statistic (for more

details, see method S6 and table S2). The similarity of the four bias rate functions indicated that the

selection of the single gene mRNA transcript datasets had little impact on modeling non-uniform read

218 distribution along mRNA transcripts, implying the universal common non-uniform read distribution of

219 different mRNA transcripts of *E. coli*. Specifically, we used the average of these four coefficients as the

final coefficients of the exponential function, which was  $f(x) = ae^{bx}$  with a = 0.256 and b = 0.256

221 **0.00128**.

# Please place Fig. 2 here.

# 223 ATUs predicted by SeqATU reach precision and recall over 0.64

224	The performance	evaluation was	conducted by	<sup>v</sup> comparing the	predicted ATUs with the ATUs in
	1		J		

225 SMRT\_M9Enrich and SMRT\_RiEnrich, which were generated based on the third-generation sequencing

and are not sensitive to transcripts with low expression levels. For a more accurate and fair evaluation,

- 227 maximal ATU clusters after pre-selection were retained in the subsequent evaluations (more details
- about the pre-selection of maximal ATU clusters can be seen in method S7 and fig. S3).

229 The precision and recall of the predicted ATUs were calculated for each maximal ATU cluster. By

230 considering only perfect matching, the average precision and recall were 0.67 and 0.67 for

231 M9Enirch\_Seq and 0.64 and 0.68 for RiEnrich\_Seq, respectively. When using relaxed matching, the

average precision and recall increased to 0.77 and 0.75 for M9Enrich\_Seq and 0.74 and 0.76 for

233 RiEnrich\_Seq, respectively. The statistics for precision and recall on maximal ATU clusters with

different sizes, as shown in Fig. 3A and fig. S4A. These results showed that the average precision and

recall were decreasing with the increasing size of maximal ATU clusters (other than several large size

236 ones due to their small number of counts). The results also indicated that the evaluation results based on

237 relaxed matching were significantly higher than those based on perfect matching across different sizes.

238 This result implied that the incorrectly predicted ATUs by SeqATU based on perfect matching tended to

have strong similarities with the ATUs in the evaluation data. In addition, we also found that more than a

240 quarter of the incorrectly predicted ATUs (25%/29% for M9Enrich Seq/RiEnrich Seq) by SeqATU

based on perfect matching matched with the transcription units in RegulonDB (19).

242	The two evaluation datasets (SMRT_M9Enrich and SMRT_RiEnrich) were both from SMRT-
243	Cappable-seq, while one of the processing steps of the technique filtered RNA reads smaller than 1,000
244	bp (6), which indicated that the ATUs in these two evaluation datasets were not comprehensive. To
245	address this issue, we enriched the evaluation data by adding the ATUs defined by SEnd-seq (7), as
246	SEnd-seq did not introduce any filtering based on RNA size. When we used the new evaluation data, the
247	ATUs predicted by SeqATU improved by 15% (0.77) and 19% (0.76) in terms of the average precision
248	based on perfect matching for M9Enrich_Seq and RiEnrich_Seq, respectively, and by 9% (0.84) and
249	12% (0.83) based on relaxed matching. The statistics for precision across different sizes of the maximal
250	ATU clusters are shown in Fig. 3B and fig. S4B, showing that the values of precision based on perfect
251	matching were significantly improved across different sizes of maximal ATU clusters by using the
252	evaluated ATUs from SMRT-Cappable-seq and SEnd-seq. This result suggested that the ATUs we
253	predicted, which were not in SMRT_M9Enrich and SMRT_RiEnrich, may be due to the RNA length
254	selection of SMRT-Cappable-seq. We enriched the evaluation data by adding the ATUs in RegulonDB
255	(19) and also found the improvement of precision across different sizes of maximal ATU clusters for
256	M9Enrich_Seq and RiEnrich_Seq (fig. S4C).
257	Furthermore, to facilitate the understanding of the performance of SeqATU and to measure the
258	influence of the maximal ATU clusters from rSeqTU on our ATU prediction method, SMRT maximal

259 ATU clusters collected from SMRT\_M9Enrich and SMRT\_RiEnrich (for more details, see method S8)

260	were applied for the CQP in two conditions (M9 minimal medium and Rich medium). We found that
261	precision and recall increased to 0.73 and 0.77 for M9Enrich_Seq, respectively, and 0.69 and 0.80 for
262	RiEnrich_Seq based on perfect matching (fig. S4D). Additionally, when using relaxed matching,
263	precision and recall significantly increased to 0.82 and 0.84 for M9Enrich_Seq, respectively, and 0.79
264	and 0.86 for RiEnrich_Seq (fig. S4D). The significantly improved results verified the ability of SeqATU
265	to accurately predict ATU when giving more accurate maximal ATU clusters. In addition, we found that
266	the number of predicted ATUs and the evaluated ATUs under the maximal ATU cluster with the same
267	size were similar except for the maximal size (Fig. 3C), and they were far less than the theoretical
268	number, which indicated that SeqATU can effectively exclude most of the incorrect ATUs.
269	Please place Fig. 3 here.
270	The bias rate constraints efficiently improve the ability of SeqATU to predict ATUs
271	We tried to use SeqATU without bias rate constraints to predict the ATUs of <i>E. coli</i> and found that its
272	performance significantly decreased compared with SeqATU (Fig. 4 and fig. S5). Specifically, the F-
273	score of SeqATU without bias rate constraints was 0.69/0.68 based on perfect matching for
274	M9Enrich_Seq/RiEnrich_Seq, compared with 0.75/0.74 for SeqATU. When using relaxed matching, the
275	
	F-score of SeqATU without bias rate constraints was 0.79/0.78 for M9Enrich_Seq/RiEnrich_Seq,
276	F-score of SeqATU without bias rate constraints was 0.79/0.78 for M9Enrich_Seq/RiEnrich_Seq, compared with 0.83/0.83 for SeqATU. This result suggested that the bias rate constraints of SeqATU

mRNA transcripts (*32-35*) and then efficiently improve the ability of the model to predict complex
ATUs.

280

# Please place Fig. 4 here.

#### 281 ATUs predicted by SeqATU display a dynamic composition and overlapping nature

A total of 2,973 distinct ATUs were identified in M9 minimal medium, and 2,767 were identified in Rich 282 medium. Among them, there were 1,423/1,550 distinct ATUs on the forward strand and 1,323/1,444 on 283 the reverse strand for M9Enrich Seq/RiEnrich Seq. Each of the predicted ATUs was comprised of an 284 average of 2.59 genes, with the largest ATU containing 28 genes across the two conditions. The 285 286 distribution of the size of the predicted ATUs is shown in Fig. 5A, from which we can see that the majority of ATUs (more than 87%) contained fewer than five genes in M9 minimal medium and Rich 287 288 medium. Approximately 41% of the genes in E. coli were contained in more than one ATU for M9Enrich Seq, compared to 43% genes for RiEnrich Seq, suggesting that the ATUs in a maximal ATU 289 290 cluster generally overlapped with each other (Fig. 5B). In addition, there were 1,576 ATU maximal clusters for M9Enrich Seq and 1,512 ATU maximal clusters for RiEnrich Seq. SeqATU identified a 291 total of 1,977 identical ATUs under the two conditions, whereas there were 1,786 distinct ATUs. Among 292 293 the distinct ATUs across the two conditions, 394 ATUs were from the same maximal ATU clusters in the two maximal ATU cluster datasets, and the rest were from different maximal ATU clusters. The fact 294 there were distinct ATUs under the two conditions suggests that ATUs are dynamically responsive to 295

different conditions or environmental stimuli (for more real examples about the ATUs under differentconditions, see fig. S6).

298	The dynamic composition of predicted ATUs by SeqATU is of great significance to understand the
299	interactions inside polymicrobial communities. For example, chronic airway infection by Pseudomonas
300	aeruginosa considerably contributes to lung tissue destruction and impairment of pulmonary function in
301	cystic-fibrosis (CF) patients (39). Marie et al. found that the presence of E. coli complemented the
302	growth defect of a <i>P. aeruginosa bioA</i> -disrupted mutant that is unable to grow on rich medium, and can
303	be beneficial to <i>P. aeruginosa</i> when biotin supply is limited (39). An ATU with a high expression level
304	coded by the <i>uvrB</i> gene is identified by SeqATU in Rich medium, while it does not exist in M9 minimal
305	medium (Fig. 6). We predicted the <i>uvrB</i> gene to be involved in the biotin metabolism pathway, as the
306	<i>bioB</i> , <i>bioF</i> , <i>bioC</i> , and <i>bioD</i> genes contained in a same ATU with it have been known in the biotin
307	metabolism KEGG pathway. Therefore, the observation by Marie et al. can be explained that the ATUs
308	coded by the <i>uvrB</i> gene of <i>E</i> . <i>coli</i> can provide the biotin supply for <i>P</i> . <i>aeruginosa</i> under rich medium.
309	This result showed that SeqATU could increase our understanding of interspecies competition and
310	cooperation, which play an important role in shaping the composition and structure of polymicrobial
311	bacterial populations.

312

Please place Fig. 5 here.

313

Please place Fig. 6 here.

# 314 Predicted ATUs by SeqATU are verified by experimental TSSs and TTSs

315	An experimental TSS dataset of <i>E. coli</i> from SEnd-seq (7) and a TF binding site dataset of <i>E. coli</i> from
316	the experimental dataset of RegulonDB (19) were used to further verify the reliability of SeqATU and
317	were named dataset 1 and dataset 2, respectively. There were 5,512 experimental TSSs in dataset 1 and
318	3,220 experimental TF binding sites in dataset 2. We considered the 5'-end genes and no 5'-end genes of
319	the predicted ATUs by SeqATU. A gene that is not the 5'-end gene of any predicted ATU is named a no
320	5'-end gene. We identified 2,177/2,005 5'-end genes and 1,266/1,160 no 5'-end genes of the predicted
321	ATUs for M9Enrich_Seq/RiEnich. A gene validated by experimental TSSs or TF binding sites means
322	that it is the immediate downstream gene of an experimental TSS or TF binding site. As a result, the
323	proportion of 5'-end genes of the predicted ATUs that were validated by experimental TSSs or TF
324	binding sites was over 1.7 times greater than that of the no 5'-end genes (Table 1). Specifically, the
325	proportion of 5'-end genes (29%/30% for M9Enrich_Seq/RiEnrich_Seq) validated by experimental TF
326	binding sites was over three times greater than the no 5'-end genes (9.2%/9.0% for
327	M9Enrich_Seq/RiEnrich_Seq). These results further verified the reliability of the ATUs predicted by
328	SeqATU in terms of the TSS level. In addition, four other experimental TSS or promoter datasets from
329	RegulonDB (19), dRNA-seq (14), and Cappable-seq (13) were also examined. The results are shown in
330	table S3, and we also found a higher proportion of 5'-end genes of the predicted ATUs validated by
331	experimental TSSs or promoters than that of no 5'-end genes.

We also used two experimental TTS datasets of *E. coli* from SEnd-seq (7) and RegulonDB (19) to

333	verify the reliability of predicted ATUs by SeqATU in terms of TTS level. These two experimental TTS
334	datasets were named dataset 3 and dataset 4, respectively. There were 1,540 experimental TTSs in
335	dataset 3 and 367 experimental TTSs in dataset 4. We considered the 3'-end genes and no 3'-end genes
336	of the predicted ATUs by SeqATU. A gene that is not the 3'-end gene of any predicted ATU is named a
337	no 3'-end gene. We identified 2,290/2,187 3'-end genes and 1,153/978 no 3'-end genes of the predicted
338	ATUs for M9Enrich_Seq/RiEnrich_Seq. A gene validated by experimental TTSs means that it is the
339	immediate upstream gene of an experimental TTS. As a result, the proportion of 3'-end genes of the
340	predicted ATUs that were validated by experimental TTSs was over two times greater than that of no 3'-
341	end genes (Table 2). Specifically, the proportion of 3'-end genes (51%/53% for
342	M9Enrich_Seq/RiEnrich_Seq) validated by experimental TTSs from SEnd-seq was over three times
343	greater than that of no 3'-end genes (15%/14% for M9Enrich_Seq/RiEnrich_Seq). These results further
344	verified the reliability of the ATUs predicted by SeqATU in terms of the TTS level. In addition, two
345	other computationally predicted TTS datasets from the works by Nadiras et al. (40) and Kingsford et al.
346	(41) were also examined. The results are shown in table S4, and we also found the proportion of 3'-end
347	genes (63%/62% for M9Enrich_Seq/RiEnrich_Seq) validated by computationally predicted Rho-
348	independent TTSs was over two times greater than that of no 3'-end genes (29%/29% for
349	M9Enrich_Seq/RiEnrich_Seq).

350

Please place Table 1 here.

Please place Table 2 here.

351

# 352 The gene pairs frequently encoded in the same ATUs are more functionally related than those that 353 can belong to two distinct ATUs

354 Functional analysis was conducted by integrating GO terms from the Gene Ontology (GO) database (42). In detail, we measured the level of functional relatedness for two types of consecutive gene pairs, 355 which is similar to the definition in the work by Mao et al. (38). Two types of consecutive gene pairs 356 were (i) gene pairs each consisting of a 5'-end gene of an ATU and the gene in its immediate upstream 357 on the same strand and (ii) all the other gene pairs inside an ATU (Fig. 7A). In addition, we used a 358 scoring scheme to measure the GO-based functional similarity between a pair of genes by Wu et al. (43). 359 This study developed a GO similarity score and showed that the larger the score, the more likely that 360 two genes are functionally related. In brief, the GO similarity score of a gene pair  $g_k$  and  $g_j$  is 361 denoted as  $S_{GO}(g_k, g_i)$ : 362

363 
$$S_{GO}(g_k, g_j) = max_{V_k \in V(g_k), V_j \in V(g_j)} s(V_k, V_j)$$

where  $V_k$  and  $V_j$  are the GO terms assigned to  $g_k$  and  $g_j$ , respectively;  $s(V_k, V_j)$  is the maximal number of common terms between paths in the two GO graphs induced by the GO terms  $V_k$  and  $V_j$ . As a result, the mean GO similarity score was higher for type-*ii* gene pairs (5.97 *versus* 4.04 for M9Enrich\_Seq and 5.86 *versus* 3.91 for RiEnrich\_Seq) than for type-*i* gene pairs. A total of 574/524 type-*ii* gene pairs had GO similarity scores greater than four (64%/63% of a total of 899/834), while only 461/404 type-*i* gene pairs had GO similarity scores greater than four (36%/34% of a total of 1,274/1,179) for M9Enrich\_Seq/RiEnrich\_Seq. We also applied a  $\chi^2$ -test (44) to determine whether the

371	distribution of $S_{GO}(g_k, g_j)$ was different for the type- <i>i</i> gene pairs and type- <i>ii</i> gene pairs. The $\chi^2$ -
372	statistics corresponded to a <i>P</i> -value less than $10^{-4}$ , which revealed that the distribution of $S_{GO}(g_k, g_j)$
373	for the type- <i>ii</i> gene pairs was significantly different from the type- <i>i</i> gene pairs. Fig. 7B shows the
374	distribution of $S_{GO}(g_k, g_j)$ for the type- <i>i</i> gene pairs and the type- <i>ii</i> gene pairs. These results strongly
375	indicated that the type- <i>ii</i> gene pairs had a higher degree of GO similarity than the type- <i>i</i> gene pairs,
376	suggesting that the gene pairs frequently encoded in the same ATUs (type- <i>ii</i> gene pairs) are more
377	functionally related than those that can belong to two distinct ATUs (type- <i>i</i> gene pairs).
378	We also carried out a similar analysis of the two different gene pairs based on KEGG enrichment
379	analysis (see more details in method S9) and found that the proportion of type- <i>ii</i> gene pairs (59%/57%
380	for M9Enrich_Seq/RiEnrich_Seq), whose two genes were contained in the same KEGG pathway, was
381	higher than the proportion of type- <i>i</i> gene pairs (32%/28% for M9Enrich_Seq/RiEnrich_Seq) (Fig. 7C).
382	The distribution of the KEGG similarity scores of the two different types of gene pairs is shown in Fig.
383	7D, suggesting that genes of type- <i>ii</i> gene pairs have a higher probability of participating in the same
384	KEGG pathway than those of type- <i>i</i> gene pairs.

385

Please place Fig. 7 here.

# 386 **DISCUSSION**

We developed SeqATU, the first computational method for genome-scale ATU prediction by analyzing
 next- and third-generation RNA-Seq data, using a CQP model. Linear constraints provided by the bias

389	rate of read distribution were, for the first time, integrated into the CQP model. Positional bias refers to
390	the non-uniform distribution of reads over different positions of a transcript (33, 35), which is handled
391	by learning non-uniform read distributions from given RNA-Seq reads (32) or modeling the RNA
392	degradation (45). The bias rate function we proposed can address the non-uniform read distribution
393	along mRNA transcripts and also be desirable for standard next-generation RNA-Seq data that involves
394	more degraded mRNAs, as the exponential function has been used to model the degradation of mRNA
395	transcripts (45). As a result, a total of 2,973 distinct ATUs for M9Enrich_Seq and 2,767 distinct ATUs
396	for RiEnrich_Seq were identified by SeqATU. The precision and recall reached 0.67/0.64 and 0.67/0.68,
397	respectively, based on perfect matching and 0.77/0.74 and 0.75/0.76, respectively, based on relaxed
398	matching for M9Enrich_Seq/RiEnrich_Seq. We further validated predicted ATUs using experimental
399	transcription factor binding sites or transcription termination sites from RegulonDB and SEnd-Seq. In
400	addition, the proportion of the 5'- or 3'-end genes of predicted ATUs that were validated by
401	experimental transcription factor binding sites and transcription termination sites was over three times
402	greater than that of no 5'- or 3'-end genes, demonstrating the high reliability of predicted ATUs. Gene
403	pairs frequently encoded in the same ATUs were more functionally related than those that can belong to
404	two distinct ATUs according to GO and KEGG enrichment analyses. These results demonstrated the
405	reliability and accuracy of our predicted ATUs, implying the ability of SeqATU to reveal the
406	transcriptional architecture of the bacterial genome.

407 In fact, the ATU architecture of bacteria is much more complex than that determined with currently

408	used experimental techniques. We investigated the 5'-end genes and no 5'-end genes of the experimental
409	ATUs identified by SMRT-Cappable-seq (6) using a combination of experimental TSSs from
410	RegulonDB (19), dRNA-seq (14), Cappable-seq (13), and SEnd-seq (7). As a result, we found that the
411	proportion of 5'-end genes (99%) validated by experimental TSSs was not significantly different from
412	that of no 5'-end genes (92%). The high percentage of no 5'-end genes validated by experimental TSSs
413	implied that the ATUs identified by experimental techniques are only a small proportion of the
414	comprehensive ATUs in bacterial organisms due to the dynamic mechanisms of ATUs. These results
415	further verified the necessity of developing robust computational methods for ATU identification.
416	SeqATU not only provides a powerful tool to understand the transcription mechanism of bacteria but
417	also provides a fundamental tool to guide the reconstruction of a genome-scale transcriptional regulatory
418	network. First, the ATU structure can help us to make new functional predictions, as genes in an ATU
419	tend to have related functions. Second, ATUs can elucidate condition-specific uses of alternative sigma
420	factors (8, 46). For example, the <i>thrLABC</i> operon is regulated by transcriptional attenuation. Totsuka <i>et</i>
421	al. found that under the log phase growth condition, the thrLABC operon is the only transcript, while
422	two transcripts are found under stationary phase growth condition, the <i>thrLABC</i> and <i>thrBC</i> . As validated
423	experimentally, $\sigma^s$ can regulate the additional promoter located in front of <i>thrB</i> under the stationary
424	phase growth condition and then separately regulate <i>thrBC</i> , which elucidates the condition-specific uses
425	of $\sigma^{S}$ (8). Third, understanding the ATU structure is of great help to construct transcriptional and
426	translation regulatory networks, such as for the construction of the $\sigma$ -TUG ( $\sigma$ -factor-transcription unit

427	gene) network (47). The transcription regulatory network consists of nodes (ATU and regulatory
428	proteins) and links (interactions) (48), and the comprehensive ATU structure can provide a nearly
429	complete set of nodes, which can improve the accuracy of regulatory prediction.
430	Although SeqATU has obtained satisfactory predicted results, there are still several challenges
431	regarding the computational prediction of ATUs. On the one hand, due to the influence of the 3'
432	untranslated region (UTR) and 5' untranslated region (UTR) in the intergenic regions, the expression
433	value of intergenic regions cannot be reproduced perfectly by the same calculation used for the
434	expression value of genetic regions. Without accurate reproduction, it is difficult to obtain the best
435	expression combination of ATUs by the programming model based on the expression value of genetic
436	and intergenic regions. On the other hand, due to the lack of strand-specific RNA-Seq data, it is difficult
437	to distinguish the expression level of intergenic regions between two consecutive genes on the same
438	strand derived from ATUs containing these two genes or antisense RNAs (asRNAs) (6, 49). All of these
439	challenges and the great significance of ATU prediction inspire and encourage us to discover more
440	information to determine the ATU structure in bacteria. For example, we plan to add high confidence
441	TSSs and TTSs information to our programming model in the future. Additionally, since the microbiome
442	is increasingly recognized as a critical component in human diseases, such as inflammatory bowel
443	disease (50), antibiotic-associated diarrhoea (51), neurological disorders (52), and cancer (53) (54),
444	predicting new ATUs of uncultured species from metagenomic and metatranscriptomic data is of great
445	significance in uncovering new regulatory pathway and metabolic products during the development of

446	diseases (55). However, due to a majority of species with unknown genomes or genome annotations
447	within a microbial community, ATU prediction on metagenomics and metatranscriptomics is still a
448	challenging task, which encourage us to pay more attention on it.

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# 584 interests: The authors declare that they have no competing interests. Data and materials availability:

- 585 The raw data and source code of SeqATU and a detailed tutorial can be found at
- 586 <u>https://github.com/OSU-BMBL/SeqATU</u>.

#### 587 FIGURES AND TABLES

- 588 Table 1. Results of predicted ATUs verified by experimental TSSs or TF binding sites. Overview of
- the experimental TSS and TF binding site datasets (dataset 1 and dataset 2) and the proportion of 5'-end
- 590 genes and no 5'-end genes of the predicted ATUs by SeqATU for M9Enrich\_Seq and RiEnrich\_Seq, which

# 591 were validated by experimental TSSs or TF binding sites.

		dataset 1	dataset 2
S	ource	In at $al$ (7)	RegulonDB TF binding
50	Jurce	Ju <i>et al</i> . (7)	sites
Tec	hnique	SEnd-seq	Collection
TSSs/TF	binding sites	5,512	3,220
M9Enrich_Se	5'-end genes	83%	29%
q	no 5'-end genes	47%	9.2%
DiFraviah Sag	5'-end genes	89%	30%
RiEnrich_Seq	no 5'-end genes	44%	9.0%

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Table 2. Results of predicted ATUs verified by experimental TTSs. Overview of the experimental
TTS datasets (dataset 3 and dataset 4) and the proportion of 3'-end genes and no 3'-end genes of the
predicted ATUs by SeqATU for M9Enrich\_Seq and RiEnrich\_Seq, which were validated by
experimental TTSs.

		dataset 3	dataset 4
Source		Ju <i>et al</i> . (7)	RegulonDB TTSs
Technique		SEnd-seq	Collection
Т	TTSs	1,540	3,67
M9Enrich_Se	3'-end genes	51%	11%
q	no 3'-end genes	15%	5.2%
D'E	3'-end genes	53%	11%
RiEnrich_Seq	no 3'-end genes	14%	4.8%

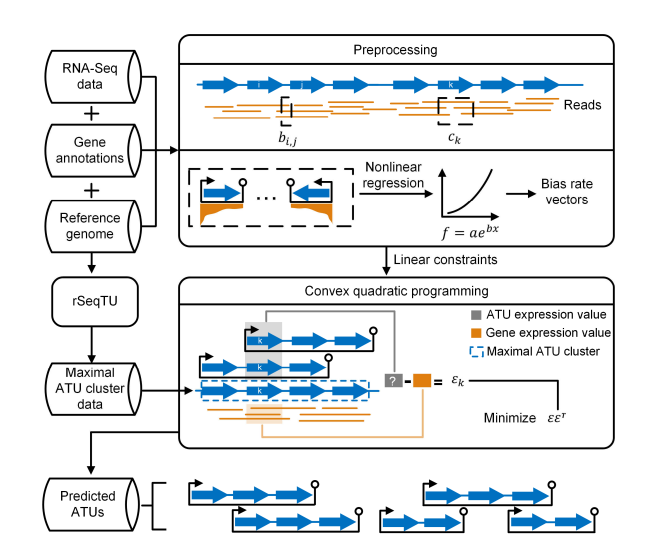
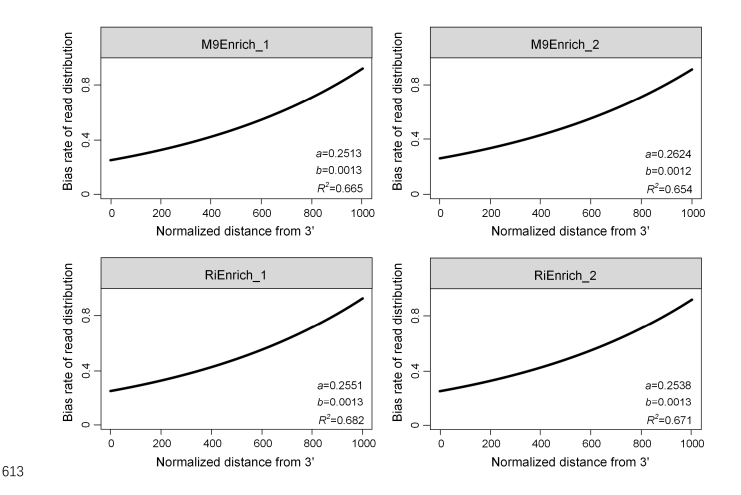


Fig. 1. Schematic overview of SeqATU. The blue arrow and orange line denote gene and RNA-Seq
 read, respectively. The preprocessing stage requires RNA-Seq data in the FASTQ format, the reference

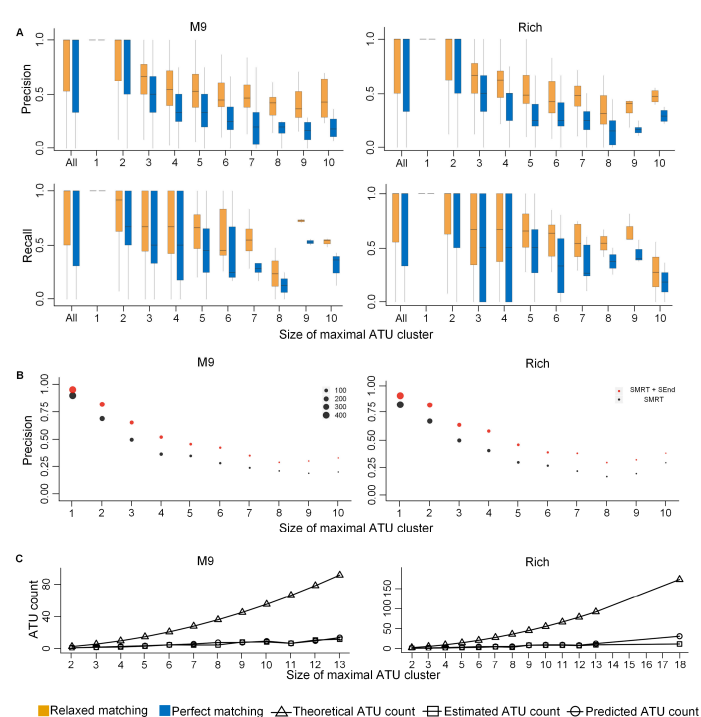
603	genome sequence in the FASTA format, and gene annotations in the GFF format, generating linear
604	constraints for the next convex quadratic programming (CQP) stage. There are two steps in the
605	preprocessing stage: (i) calculating the expression value of the genetic region $c_k$ and intergenic region
606	$b_{i,j}$ and ( <i>ii</i> ) modelling non-uniform read distribution along mRNA transcripts; specifically, we acquired
607	a bias rate function $f(x) = ae^x$ using nonlinear regression and then constructed genetic or intergenic
608	region bias rate vectors. The maximal ATU cluster data determined by rSeqTU and the linear constraints
609	from preprocessing are both taken as inputs of CQP. CQP seeks the optimum expression combination of
610	all of the to-be-identified ATUs to minimize the gap $\varepsilon \varepsilon^T$ between the predicted ATU expression profile
611	and the genetic and intergenic region expression profile. Finally, the output of CQP is the predicted
612	ATUs.



614 Fig. 2. Results of modelling non-uniform read distribution along mRNA transcripts. The four bias

rate functions  $(y = ae^{bx})$  by nonlinear regression had similar coefficients (a and b) across the four datasets M9Enrich 1, M9Enrich 2, RiEnrich 1 and RiEnrich 2.

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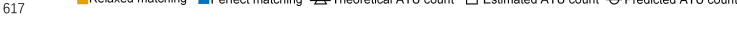


Fig. 3. Overall evaluation results of SeqATU. (A) Precision and recall based on perfect matching and

619 relaxed matching for M9Enrich Seq (left) and RiEnrich Seq (right) using evaluated ATUs from SMRT-

- 620 Cappable-seq. (B) Average precision based on perfect matching for M9Enrich Seq (left) and
- 621 RiEnrich\_Seq (right) using evaluated ATUs from SMRT-Cappable-seq (black) and evaluated ATUs from
- 622 SMRT-Cappable-seq and SEnd-seq (red). The magnitude of the point denotes the number of maximal
- 623 ATU clusters with same size. (C) Average number of ATUs across different sizes of SMRT maximal
- 624 ATU clusters for M9Enrich\_Seq (left) and RiEnrich\_Seq (right).

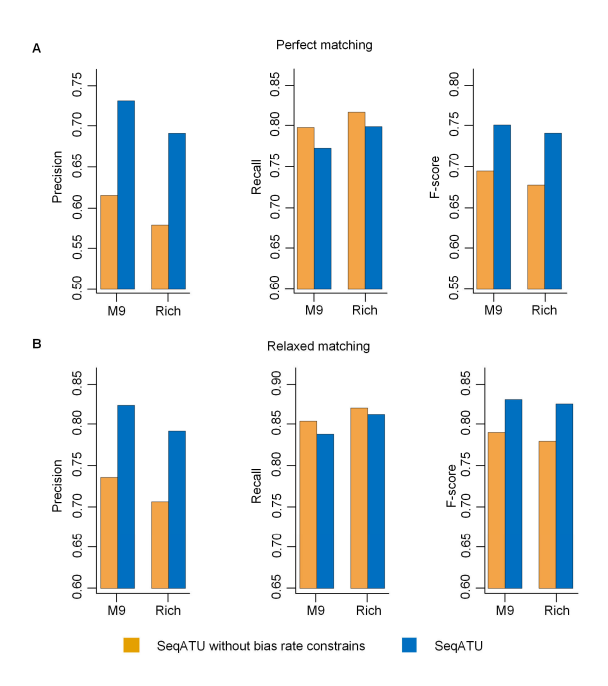
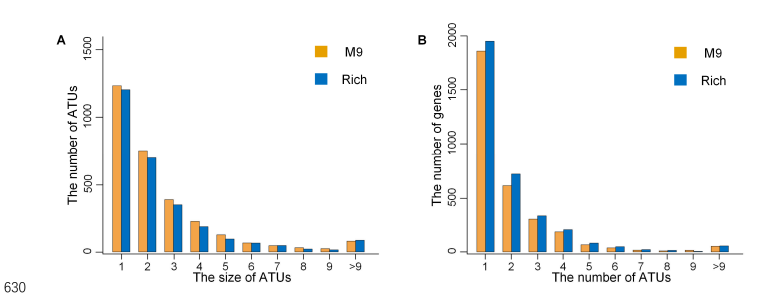


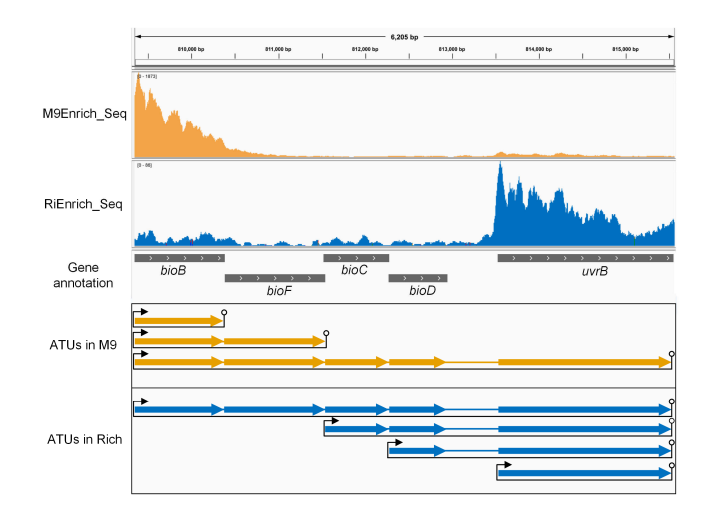
Fig. 4. Comparative analysis of the performance between SeqATU and SeqATU without the bias
rate constrains for SMRT maximal ATU clusters. (A) Precision, recall and F-score based on perfect
matching for M9Enrich\_Seq and RiEnrich\_Seq. (B) Precision, recall and F-score based on relaxed
matching for M9Enrich\_Seq and RiEnrich\_Seq.



631 Fig. 5. Comprehensive analysis of the predicted ATUs by SeqATU. (A) Number of ATUs across

632 different sizes. The size of an ATU is the number of its component genes. (B) Distribution of the number

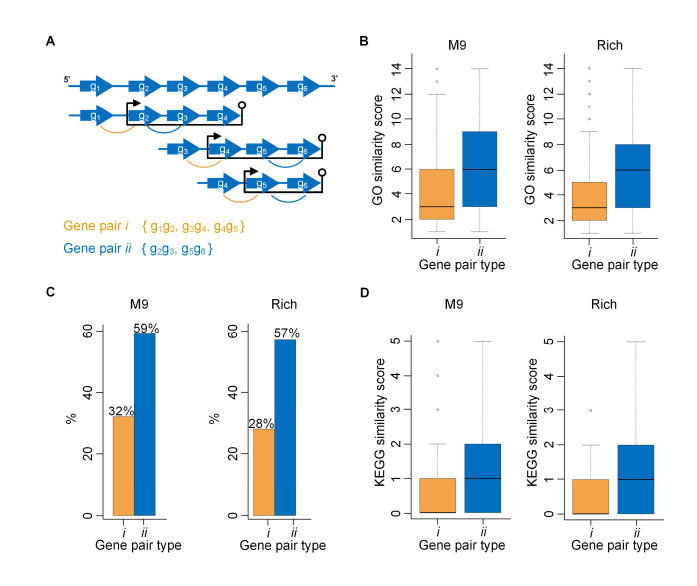
633 of ATUs per gene.



# 635 Fig. 6. Integrative Genomics Viewer (IGV) representation of the mapping and ATUs. Mapping and

636 ATUs of M9Enrich\_Seq (orange) and RiEnrich\_Seq (blue) were shown for the maximal ATU cluster

<sup>637</sup> containing the *bioB*, *bioF*, *bioC*, *bioD* and *uvrB* genes.



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