

1 Title

2 Strategies for building computing skills to support microbiome analysis: a five-year perspective
3 from the EDAMAME workshop

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17
18 Abstract

19 Here, we report our educational approach and learner evaluations of the first five years of the
20 Explorations in Data Analysis for Metagenomic Advances in Microbial Ecology (EDAMAME)
21 workshop, held annually at Michigan State University's Kellogg Biological Station from 2014-
22 2018. We hope this information will be useful for others who want to organize computing-
23 intensive workshops and encourage quantitative skill development among microbiologists.

24

25 Importance

26 High-throughput sequencing and related statistical and bioinformatic analyses have become
27 routine in microbiology in the past decade, but there are few formal training opportunities to
28 develop these skills. A week-long workshop can offer sufficient time for novices to become
29 introduced to best computing practices and common workflows in sequence analysis. We
30 report our experiences in executing such a workshop targeted to professional learners
31 (graduate students, post-doctoral scientists, faculty, and research staff).

32

33 Introduction

34 It is now recognized that microbial communities (“microbiomes”) play essential roles for
35 the health of the environments and the hosts that they inhabit. In addition, advances in high-
36 throughput sequencing technologies allow for observations of the diversity and functional
37 potential of microbiomes in their habitats (1), captured with spatially and temporally ambitious
38 study designs (2). Together, these advances in knowledge and methodology deepen and
39 broaden our understanding of the centrality of microbiomes for host and environmental health.
40 Because of the economy and accessibility of high-throughput sequencing, researchers can now
41 investigate the diversity of interesting microbiomes and can begin to untangle how this
42 diversity contributes to host or ecosystem health. Efforts to capitalize on the promise of
43 microbiome sequencing data have resulted in information-rich genomic datasets that must be
44 analyzed to gain knowledge of their intricate relationships.

We realized that there was a need for broad computational training in microbiome analysis. In 2014, we were encouraged by Dr. C. Titus Brown (now at University of California-Davis) to offer a microbiome analysis workshop. At the time, he led the Analyzing Next-Gen Sequencing (ANGUS, <https://angus.readthedocs.io/en/2018/index.html>) Workshop at Michigan State University's Kellogg Biological Station (KBS). He noted that some ANGUS learners were particularly interested in microbiome analysis and that there were limited offerings for this training. At the time, there were several short-duration workshops focused on specific tools, such as QIIME(4) and mothur(5), as well as a broader, multi-week course, STAMPS (<https://www.mbl.edu/education/courses/stamps/>), at the Marine Biological Laboratory in Woods Hole, MA USA. There were few workshops that addressed the needs of learners who wanted more information than what could be covered in a day but also could not commit to spending several weeks away. Thus, we suspected that there was a need for broad and economical training in microbiome analysis, especially in the U.S. Midwest.

In response, we created a one-week intensive course to train biologists (from graduate students to faculty) in microbiome-associated sequencing analysis, from raw sequence handling and quality control to statistical analyses and experimental design. We named the course EDAMAME: Explorations in Data Analysis for Metagenomic Advances in Microbial Ecology. Ashley Shade, at the time a new assistant professor in microbial ecology at the Department of Microbiology and Molecular Genetics at Michigan State University, initiated the workshop and started its content development from her materials from a short workshop she offered while training in her post-doctoral advisor's lab. Tracy Teal was recruited and brought her array of experience and perspective as a leader in the Software and Data Carpentries workshops, which

provide general computing training. In the first year, J. Herr, a post-doc in Shannon Manning's lab at Michigan State who had Data Carpentry training, contributed to developing and implementing the original content. The instruction team expanded in 2016 to include Adina Howe, who was a new faculty at Iowa State and brought important expertise in untargeted metagenome analysis.

Here, we report a five-year perspective on the EDAMAME workshop. We present EDAMAME's learning objectives, target audience and admissions, instructional team, learning environment, educational strategy and assessment, and community resources. We discuss results from assessment, lessons learned and an outlook for future microbiome training.

Results

EDAMAME learning objectives

EDAMAME's learning objectives were tailored annually to incorporate learners' changing interests and changes in tools and technology (**Figure 1**). As a consequence, though the overall content was not drastically changed, we created and retired tutorials as demands changed. For example, when we found that many of our learners had exposure to and experience with amplicon sequence analysis, we shortened that content to accommodate more time for metagenome analysis. However, each year featured foundational tutorials in computing literacy, state-of-the-science tools for microbiome analyses, ecological statistics, and computing best practices. We provided specific datasets to accompany each of the hands-on tutorials and

encouraged learners to use these datasets in the classroom. Some datasets were used for more than one tutorial to provide continuity through the analysis workflow.

Target learners and admissions

We targeted our applicant pool towards learners who would benefit most from the training and who we expected would share their developed expertise with others to maximize the reach of the workshop's training. We accepted applicants who were novice in their analysis skillset and did not have apparent access to other resources to support their skill development. We also aimed to promote diversity in scientific discipline (e.g., human, agricultural, environmental), learner gender and background, research institution (e.g., undergraduate-serving, research university, agencies), geography (with special advertising to learners from the Midwest), and academic level (**Figure 2, Figure 3**). We also strove to provide opportunity to international learners and learners from underrepresented backgrounds. To advertise the course, we used social media (Twitter), our website, and professional networks. We also attempted to reach broader audiences by advertising with international scientific networks, especially Ciencia Puerto Rico in 2014 - 2016.

In each workshop, we could accommodate 23 - 26 learners in the classroom, and applications were oversubscribed every year (**Table 1**). As admissions became increasingly competitive, we began to require (rather than to encourage) that applicants had generated a microbiome dataset prior to the workshop. We found that students who had struggled in an analysis attempt were highly incentivized to maximize their time at the workshop. Also, they could work on their data during office hours and ask specific questions to the instructors and Teaching Assistants (TAs).

109 *Instructional team*

110 A large instructional team was necessary to support EDAMAME’s learning goals. There
111 were one to three lead instructors per year (**Table 1**). The instructors led the course, oversaw
112 admissions, provided lectures and course content, determined guest lectures, and mentored
113 TAs in tutorial development. In the final two years of the workshop, there was also a course
114 coordinator who oversaw conference logistics, fielded learner and applicant questions, and
115 coordinated transportation for learners, guest lecturers, and instructors.

116 The hands-on nature of the workshop necessitated several dedicated TAs. Multiple
117 instructors and supportive TAs in the classroom allowed us to be immediately responsive to the
118 needs of the learners. TAs led tutorials based on interest and expertise. Having multiple TAs
119 broadened instructional expertise and allowed unscheduled time for each TA to rest when they
120 were not supporting instruction. Most often, new learners struggled with basic syntax and
121 interpreting error messages. Novice TAs (e.g., early graduate students) helped learners trouble
122 shoot common errors, while the more senior TAs and instructors assisted with more
123 complicated hurdles (e.g., software and operating system incompatibilities, experimental
124 design power for data analysis). In addition to instruction, TAs supported the logistical aspects
125 of the course, such as local transportation for learners, purchasing supplies, and assisting
126 learners with unexpected personal needs (e.g., trip to the medical center, forgotten
127 toothbrush). TAs included volunteers (graduate students and post-docs) and graduate
128 assistants partially supported by EDAMAME external funding. Participation in the workshop
129 also offered TAs benefits to engage in teaching opportunities that served diverse audiences.

There were also numerous invited guest instructors who offered tutorials, technical lectures, and research talks (**Table 2**). These guest instructor were invited to include diverse career stages and expertise across microbiome science (**Table 2**). Guest instructors varied according to guest availability, learner interests, and workshop duration, but some guest instructors generously provided content every year. Stuart Jones (University of Notre Dame) taught statistical analysis in R; Patrick Schloss and members of his lab (University of Michigan) taught amplicon analysis with mothur; Jim Tiedje (Michigan State University) provided a lecture and discussion on the future of microbial ecology. Instructors interacted with the learners during dinner and social time, and this provided an opportunity for learner networking and discussions.

Learning environment and daily schedule

EDAMAME was held at the Kellogg Biological Station (KBS), which offered a remote location, offering an immersive experience for learners and instructors. KBS was also chosen for economy – the room and board rates at KBS were affordable to many (e.g., ~\$370 per week in 2018). Teaching assistants and volunteers provided transportation from the Kalamazoo and Lansing airports to KBS. KBS also provided conference services, dining, wifi, and bonfires. Finally, the natural setting and outdoor activities at KBS provided a respite to time spent in front of the computer.

The length of the workshop varied from 7 - 11 days (**Table 1, Figure 4**), including travel days. The morning schedule included an overview lecture followed by hands-on tutorials and

group learning activities. After lunch, we had an afternoon lecture and additional tutorials. We held optional office hours with “choose your own adventure” tutorials and/or lectures on learner-chosen topics during the afternoon break. For example, in 2018 we discussed exact sequence variant analysis because it was a new approach being used in the field for defining operational taxonomic units. Learners could also ask specific questions about their own data during office hours. After dinner, we held an evening guest lecture in microbiome research. Evenings provided free time for networking and relaxation.

EDAMAME educational strategy and assessment

EDAMAME’s educational strategy addressed two training needs. First, we offered general training in the fundamentals of introductory computing (*e.g.*, command line, scripting, cloud computing, bioinformatic workflows). This equipped participants with the basic skills needed to independently execute their analyses. We also offered specific training to overcome hurdles particular to microbial metagenomic data analysis and advised on best practices for microbiome analysis. To iteratively assess these strategies, we used a combination of summative and formative assessments to determine participant learning gains.

For the summative assessments, we worked with educational consultants to develop online, anonymous surveys and perform pre- and post-workshop assessments. These assessments evaluated student-reported learning gains and confidence in areas aligned with our learning objectives. The learners created a password to preserve their anonymity while allowing for linking the pre- and post-survey responses. To maximize response rate, we

provided dedicated time in the classroom to complete the surveys. The pre-assessment survey was completed on the first full day, and the post-assessment survey was completed on the final day of the workshop. We updated the survey annually to reflect any new or changed learning objectives but maintained the structure to facilitate interannual comparisons. Results of the annual surveys guided the continued development of course materials and topics covered. In the early years of the workshop, we had consultants perform in-classroom observations and provide feedback to the instructors. Ultimately, we compiled the five years of pre- and post-survey data and performed a longitudinal analysis.

In the pre- and post- surveys, learners were asked to indicate the extent to which they understood specific learning outcomes or skills covered in the course, with ratings (e.g. Strongly Disagree, Disagree, Agree, and Strongly Agree (**Figure 5**)).

We also used “real-time” assessment during the workshop by replicating formative assessment strategies found to be effective in Software Carpentry workshops (8–10). Formative assessment is an approach where teachers use informal practices to assess student learning during the workshop in order to evaluate understanding and modify teaching if needed. We used red and green sticky notes and ‘minute cards’ to get this real-time feedback from students. Each participant was given a green (“I’m doing okay”) and a red (“I have a question”) sticky note to stick onto their open laptop during tutorials. This visual cue allowed instructors to quickly survey the classroom and determine learners’ comfort level, and to attend to any student who was struggling during tutorials. Furthermore, it allowed students to continue working through tutorials or troubleshooting without the need of raising their hand. We also

193 employed “minute cards”. After each tutorial, students wrote what went well on the green
194 sticky note and what could be improved on the red sticky note. Instructors and TAs read
195 through notes during breaks to quickly identify gaps in understanding. This allowed us to
196 identify gaps and make adjustments (e.g., in speed) in the subsequent instruction period.

197

198 *Building community resources and peer networks*

199 We were dedicated to promote a welcoming and supportive learning environment. We
200 presented a Code of Conduct in the welcome lecture, which outlines expectations for student
201 and faculty behavior during the workshop and reporting procedures if those behavior
202 expectations are not met, so that it was clear that any questionable conduct was grounds for
203 dismissal. We created an online shared document etherpad for collaborative note taking to
204 maximize engagement and inclusivity. All materials were regularly updated and available
205 online through our course website. We did our best to accommodate learners with families,
206 providing private housing to families and learners with special requirements.

207 We aimed to build a peer learning community and to provide resources to support
208 learners beyond the workshop. We offered an informal meet-and-greet on the arrival travel day
209 and get-to-know-you short talks as lightning presentations after the first full day. These
210 interactions allowed learners to identify peers with common research interests early in the
211 workshop. We created a workshop website and public repository on GitHub so that learners
212 (and outside parties) could access EDAMAME learning materials. Linked content included
213 lectures, hands-on tutorials, and reference lists. These materials have been shared openly, with

most content licensed as Creative Commons generic (e.g., CC-BY), so all course registrants and anyone else could have access. We also shared group email lists and encouraged social media outreach via Twitter and blogging. An EDAMAME meet-up was also held at the International Society for Microbial Ecology 2016 meeting in Montreal, CA.

Pre- and post-survey comparisons and qualitative interviews

Ninety-seven percent of EDAMAME learners from 2014 to 2018 rated the workshop overall in the top evaluative categories, “good” to “very good.” (**Figure 6**). A comparison of pre- and post-assessment learner-reported learning gains and/or confidence with the major learning objectives of EDAMAME show gains in all sub-categories of learning reported (**Figure 7**). There were largest gains between the pre- and post-assessments with Computational Understanding (**Figure 7B**) and Perception in Ability (**Figure 7C**).

We also asked short-answer questions at the end of the survey, in which learners were asked to design an experiment and report how they would process and analyze microbial community high-throughput sequencing data. We observed increased sophistication in the responses to the short-answer questions from the pre- to post-survey, with some learners leaving the questions blank in the pre- survey and then providing thorough answers in the post-survey (data not shown, but anonymized annual assessment reports available upon request). This suggests large gains especially for learners who were new to high-throughput sequence analysis.

Qualitative interviews from 9 learners who attended EDAMAME from 2014-2016 (each spending 25-40 minutes with the interviewer, **Table 3**) suggested that this group of learners were largely satisfied with the workshop and appreciated the attentiveness of the TAs and instructors as well as the red/green sticky note mechanism for soliciting help in real time. However, some of these learners also felt that there was too much material covered in the workshop and reported that they struggled to keep up with the pace of the course (“Content overwhelm”). Finally, we had many interviewed learners state that the workshop and materials covered made a positive impact on their career and research.

Discussion and Lessons Learned

We offer suggestions from our experiences for running an effective microbiome analysis workshop (**Figure 8**). EDAMAME’s content changed from 2014 to 2018 to meet changing learner needs. These changes were guided in part by the applicants’ responses to questions about their dataset and their expectations for the workshop. For example, amplicon analysis (e.g., 16S rRNA gene sequencing) was favored in early years while untargeted metagenome analysis was favored in later years. Similarly, proportionally fewer students in 2018 were novice to the command line or R, but the majority of the class appreciated the refresher. Some of the learners with self-taught experience embraced the opportunity to re-learn the “correct” approaches and to gain missing foundational knowledge. Several tutorials were popular every year. For example, there was a consistent demand for ecological statistics and “supporting” skills like GitHub/version control, and cloud computing.

High instructor to learner ratio was essential for the success of the hands-on EDAMAME workshop. In the years that we had the lowest instructor to learner ratios (e.g., in 2014 and 2015, **Table 1**), the TAs and instructors anecdotally reported exhaustion while the learners craved more attention. In addition to formal instructors, learners could assist one another. To facilitate peer learning, we arranged the classroom in tables with groups of two or four. We also encouraged learners to support one another with troubleshooting in the time that it would take for a free instructor to come to assist

Regardless of the length of the course, several learners indicated in their post-assessments that more time at the workshop was needed each year. However, learners who were faculty or staff researchers shared (in informal conversations) that they would have been unable to commit to a longer workshop due to other professional responsibilities. We noted that there were other offerings for multi-week workshops e.g., STAMPS), as well as several one- or two-day workshops at professional society meetings and pipeline-specific training (e.g., mothur and QIIME).

Timing the workshop had several challenges. EDAMAME was held in the summer, and we tried to avoid scheduling it for the same week as major microbiology conferences, like the American Society for Microbiology Microbe meeting, the International Symposium on Microbial Ecology (ISME) and Ecological Society of America meetings. Because microbiome analysis spans multiple disciplines, it was hard to avoid all of the large conferences that microbiome researchers may attend. We also had to change the timing workshop every year to accommodate the KBS event schedule. As EDAMAME grew in popularity, some learners applied

for fellowships or travel awards to support their training, but the annual change in timing made it difficult for students to plan. Moving the workshop to a dedicated conference site (e.g., a hotel) may help with consistent timing, but it would also increase the cost to learners.

We found that using cloud computing streamlined course content and democratized access. We used the Amazon Elastic Compute Cloud (EC2), which was cost effective and available to students who do not have access to high performance computers at their home institutions. In early years, we guided learners through software installation on the EC2, but in later years, we installed the software on the EC2 for the learners so that they could focus on moving data to and from the EC2. Using the EC2 presented a challenge for learners who were affiliated with government agencies or research laboratories (e.g., US Environmental Protection Agency , US Geological Survey) because of their need for additional security and management approval prior to installing new software or moving data. While we did not have a perfect solution for these learners, we began to anticipate their needs and prompted them in advance to receive required permissions. Another hurdle with using the EC2 was the changing way that Amazon provided student or educational computing resources over the years. In some years, Amazon provided individual credits to learners and in others required the instructors to apply for an educational grant. Cloud computing logistics needed to be anticipated about nine months in advance, but in years where individual email addresses were needed, it was impossible to prepare until after admissions were finalized, which typically occurred 4 - 6 months in advance of the workshop. We also note an issue for some international learners who did not have credit cards compatible with Amazon requirements to enroll for an EC2 account, and for these learners we had to share our own accounts or create accounts for them.

296 While our applicant pool and learner demographics reflected balance in gender, discipline,
297 and academic level, EDAMAME fell short of its racial diversity goals. We could have benefitted
298 from improvement in advertising the course to reach a broader pool to attract more applicants
299 of color. We largely advertised on social media and through word of mouth. We recommended
300 to specifically advertise to key target learner groups, like those underrepresented in the
301 sciences who may be expected to have less access to the training. On a positive note, we have
302 evidence that EDAMAME was reaching socioeconomic diversity goals, as two interview
303 respondents were clear that they would not have had the same opportunity for training and
304 advancement given their lower income backgrounds if it had not been for EDAMAME.

305 A final lesson to share is the balance between course value and learner costs. In its first
306 years, EDAMAME was funded piece-meal by generous sponsors. We experimented with a
307 mixed enrollment model of offering EDAMAME for university credit to local students and for
308 fee to outside students, but many of the local students could not afford the summer tuition
309 required for the credit hours. Then, EDAMAME was funded by external federal grants (NIH
310 2015-2018 and USDA 2017-2018.) and we stopped offering it for credit for logistical simplicity.
311 We began to charge modest workshop fees (\$325) to support items that could not be covered
312 by the grant (e.g., coffee, snacks). As soon as we began to charge workshop fees, the majority
313 of applicants began to request financial aid. We realized that many of the learners, mostly
314 graduate students and post-doc, were paying for the workshop personally, so we then worked
315 to waive fees for eligible students in need and offer scholarships for students with international
316 travel. By contrast, the instructional team did not have enough funds to fully pay the TAs and
317 instructors, who largely volunteered their time because they believed in the mission of the

318 training. Guest instructors and lecturers generously volunteered their time as part of their
319 broader impacts for federal grants, and the workshop covered their travel expenses along with
320 room and board at KBS. Thus, there is inevitable tension balancing instructor compensation and
321 course affordability.

322 How much does it actually cost to run a workshop like EDAMAME? The first year, we ran
323 the workshop for less than \$14,000; students paid their own expenses of room and board; and
324 no workshop fees were charged. This face amount did not include substantial additional
325 support that was provided via shared logistics with the ANGUS workshop, which was occurring
326 at the same time at Kellogg Biological Station. It also did not include any support for personnel,
327 which was the largest expense. Ideally, there would have been an annual budget for instructor
328 and TA summer salaries, a logistics coordinator salary, and hourly salary for undergraduate
329 labor during the course. We also realized that unless we could procure funds to support
330 personnel, the training may not be valued as highly by institutions and peers, and may instead
331 be perceived as a cost to other scholarly activities. The second biggest expense was be financial
332 aid to offset costs of room and board and workshop fees to learners who needed it, which we
333 provided in 2017 and 2018 to qualified learners, with USDA support. The third biggest expense
334 was the educational consultant to evaluate the course as a neutral third-party, which was
335 \$5,500 to \$6,000 per evaluation. The remaining expenses were conference services at Kellogg
336 Biological Station, and lodging and travel expenses for the instructional team and guest
337 speakers. In summary, there is a trade-off between the course cost, inclusive of the real value
338 of instructor/TA time, and workshop affordability for the learners.

339

340 Future directions

341 While the data indicate that EDAMAME workshop was effective, a limited number of
342 learners can be accommodated per year, and there is high effort from the instructional team to
343 support them. This is a low-throughput model of skill development. We are eager to reach a
344 larger learner pool than what we could accommodate in the classroom. In 2016, we
345 experimented with live engagement of three to five remote learners (varied by tutorial) using
346 free conference calling and screen sharing resources. The remote learners participated as a
347 group at the same location. They engaged with the lectures and tutorials as fully as possible
348 (but missed out on the guest lectures and other events). This added a mild distraction for the
349 on-site learners, but the workshop proceeded relatively smoothly. The biggest hurdle was
350 engaging with the remote learners during tutorials, as they had no classroom support. It is
351 possible that a remote learning workshop could be successful, given an appropriate investment
352 into conference technology, an on-site coordinator dedicated to its logistics, and an enhanced
353 instructional team with traveling TAs dedicated to the remote classrooms.

354 The content of EDAMAME remains freely available online
355 (<http://www.edamamecourse.org/materials/>), but parts of the content are also being
356 transitioned to local offerings. Many universities desire more offerings of online or digitized
357 curriculum, and there is a question of how to balance the university's need to provide quality
358 instruction for tuition with the open-science philosophy of providing free, democratic access to
359 information. At Michigan State University, we are developing a graduate-level learning module

on microbial metagenomics that includes amplicon and untargeted metagenome analysis pipelines. The 1-credit metagenomics module includes hands-on tutorials, is offered twice a week for one month and is accompanied by pre-recorded lectures. Post-doctoral trainees or faculty can enroll for a modest fee. Though based on EDAMAME materials, the modular content at Michigan State covers less content because there are prerequisite modules required for enrollment. Learners already have familiarity with the command line, with submitting jobs to the high-performance computing cluster, and with fundamentals of microbial genome analysis. EDAMAME materials have also been extended to international workshops including, a metagenomics one-day crash course in Rio, Brazil and a one-week microbiome analysis workshop at Centro de Investigaciones Biológicas del Noroeste in La Paz, Mexico. In addition, more general tutorials (e.g., shell, GitHub, etc) remain available from other data science training efforts, including Software and Data Carpentry, and short format 2-day workshops are available through The Carpentries (<http://carpentries.org>) on these skills.

Finally, we seek to maximize the impact of EDAMAME by offering this kind of training to those who need it most. We hope that the impact of our trainees training others is a lasting legacy of EDAMAME. We have found that our international learners have benefited immensely from this course, as they are challenged by access to compute resources or training. Going forward, we hope to continue to identify target audiences who could both benefit from our training and extend its impact broadly. Additionally, sequence analysis will continue to evolve with technologies, impacting the depth and breadth of scientific questions and experiments that are imaginable. We hope that our course content can continue to remove obstacles for scientists who wish to engage in these technologies.

382

383 Materials and Methods

384 This research was exempt under IRB ID# i052533 (standard educational practices), as
385 reviewed by the Michigan State University Biomedical, Health Sciences Institutional Review
386 Board (BIRB) and Social Science, Behavioral, Education Institutional Review Board (SIRB).

387 Data analysis for the pre- and post-survey assessment and associated reports were
388 generated by an outside research consultants. Final reports for the years 2016, 2017 and 2018,
389 were written by Beth M. Duckles, PhD of Insightful, LLC and for years 2014 and 2015, reports
390 were written by Julie Libarkin of STEM ED. LLC. Code is available at
391 [<https://github.com/ShadeLab/EDAMAMESurveys>]. Beth M. Duckles of Insightful also
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393

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Conflict of Interest

Amazon EC2 provided compute resources to EDAMAME students. In 2014, Illumina provided pizza dinner and in 2014-2017 MO BIO provided t-shirts and blogging opportunities on their company's blog.

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Figure legends

Figure 1. Overview of learning objectives for the EDAMAME workshop.

Figure 2. Distributions of EDAMAME learner gender and age, 2014-2018. Data were collected and summarized from pre- and post-workshop assessments.

Figure 3. Distributions of EDAMAME learner ethnicity and academic status, 2014-2018. Data were collected and summarized from pre- and post-workshop assessments.

Figure 4. An example of an ideal one-week schedule for EDAMAME. For a full schedule with links to 2018 EDAMAME tutorials and talks, see <https://github.com/edamame-course/2018-Tutorials/wiki/Schedule-EDAMAME-2018>. Content for all five years of the workshop are also available at <http://www.edamamecourse.org/materials/>.

Figure 5. Representative survey questions for the “Computational Understanding” scale.

Figure 6. Overall EDAMAME assessment 2014-2018. Data were collected and analyzed from pre- and post-workshop assessments.

Figure 7. Summarized comparison of self-reported learning gains between pre- and post-workshop assessments, aggregated over 2014-2018. (A) Comfort with computational tasks; (B) Computational Understanding; (C) Perception in Computing Ability; and (D) Coding Ability.

479 **Figure 8.** Lessons learned over five years of the EDAMAME workshop.

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483 Tables

484 **Table 1. Summary of EDAMAME dates, instructional staff, applicants, and learners from**
485 **2014-2018.**

Year	Dates	No. Days	No. TAs	No. Instructors	No. applicants	No. workshop learners ¹
2014	22 June to 29 June	8	1	3	50	23
2015	21 June to 01 July	11	6 ²	1	93	32 ³
2016	10 July to 20 July	11	6	3	62	25
2017	06 August to 12 August	7	7	3	63	26
2018	24 June to 30 June	7	10	2	103	26

486 ¹No. workshop learners are from pre- and post- survey responses. Additional local learners
487 participated *ad hoc* and may not have completed surveys.² There were two guest TAs in 2015
488 who participated only in one tutorial each, with the remaining 4 TAs available throughout the
489 workshop. ³2016 participant data included 3 remote learners who participated in select
490 tutorials.

491

Develop working proficiency at the command line and with shell.

Explain the process of high-throughput sequencing, provide an overview of data-handling (quality control, pre-treatment), and discuss their biases.

Access computing resources: Transfer data and run analyses on Amazon EC2 and/or a high-performance computing cluster.

Access and/or create version-controlled code and resources on GitHub.

Discuss steps in the ecological analyses of microbiomes, including alpha and beta-diversity, ordinations, and resemblance metrics.

Explore datasets and statistically test hypotheses in R.

Visualize patterns in microbial communities using R.

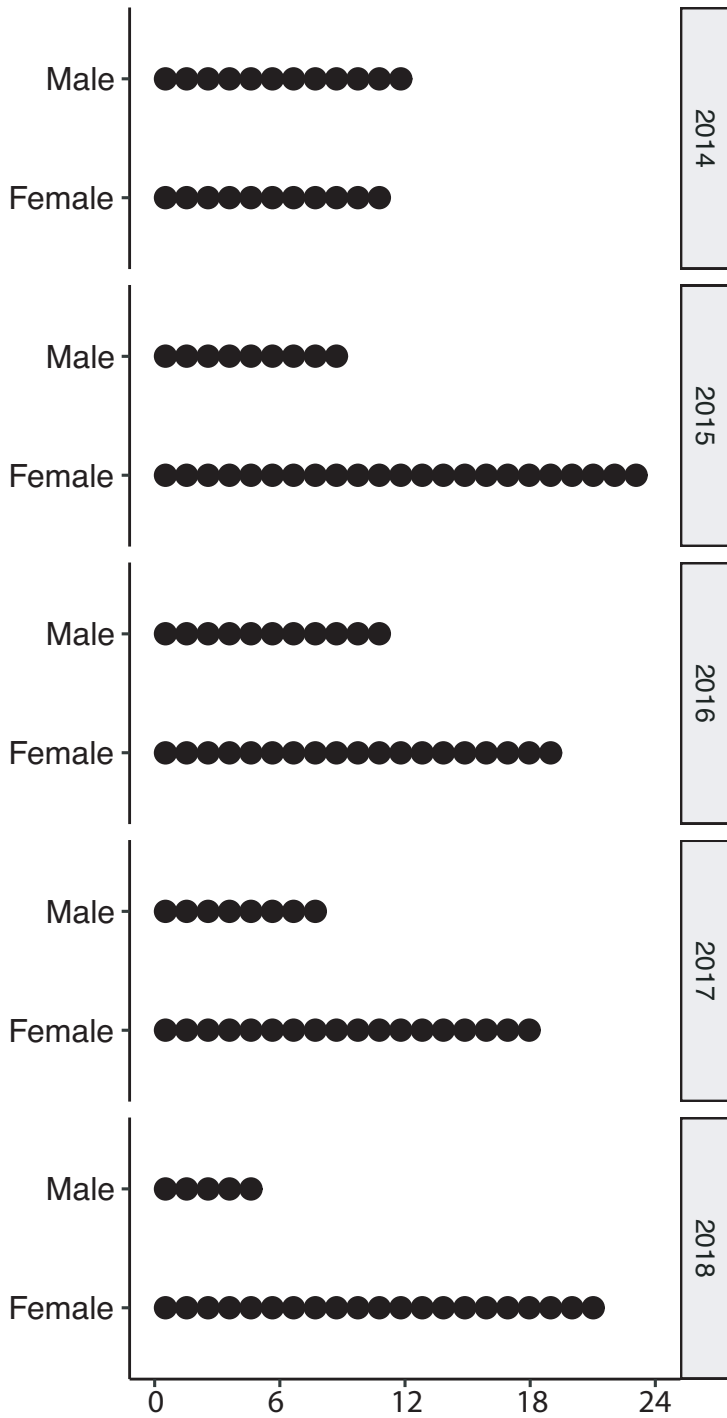
Develop a working proficiency with amplicon sequencing workflows and tools (e.g., QIIME, or mothur, or usearch).

Develop a working proficiency with shotgun metagenomics workflows.

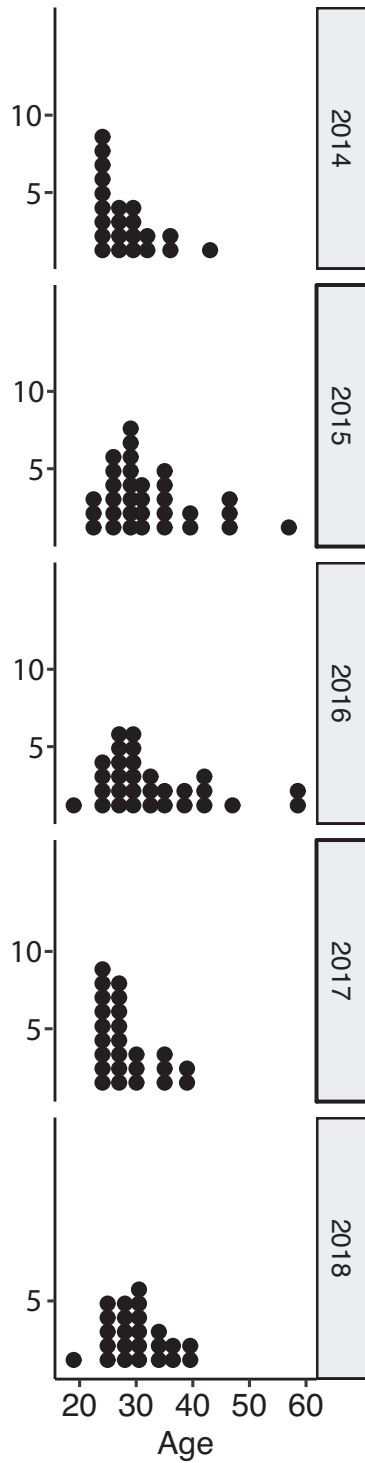
Become familiar with publicly accessible microbial sequence databases/repositories (e.g., NCBI, MG-RAST, FunGene) and the tools that they offer for deposition and analyses.

Identify resources for troubleshooting. This includes: how to ask for and where to find general help online, through peer networks, and from workflow-specific resources (e.g., public tutorials and wikis).

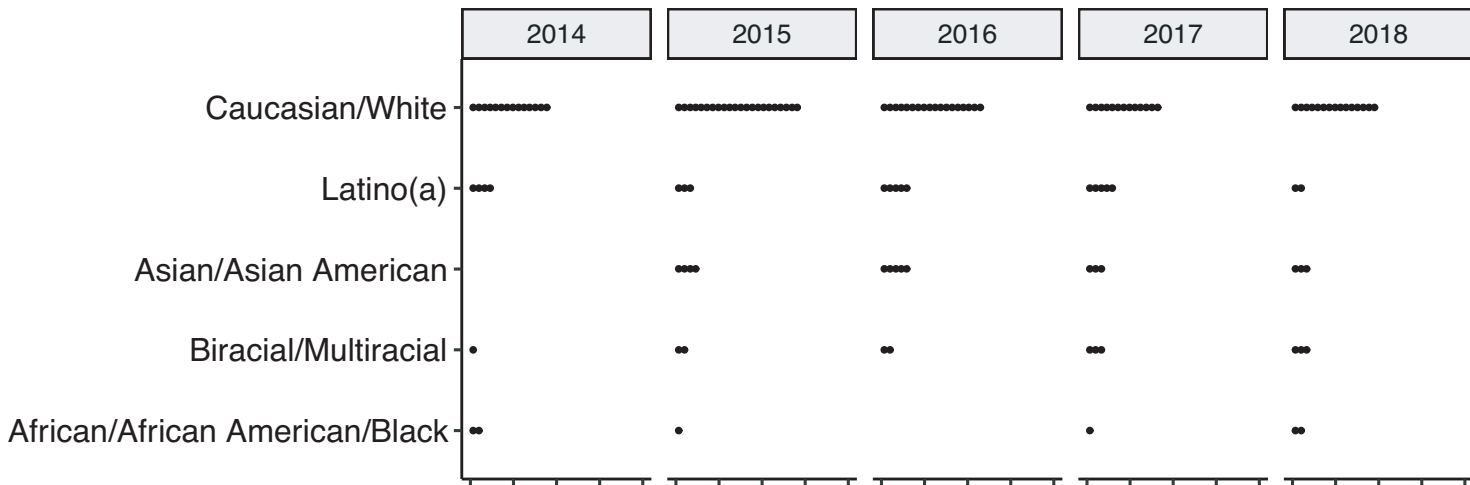
Attendee Gender



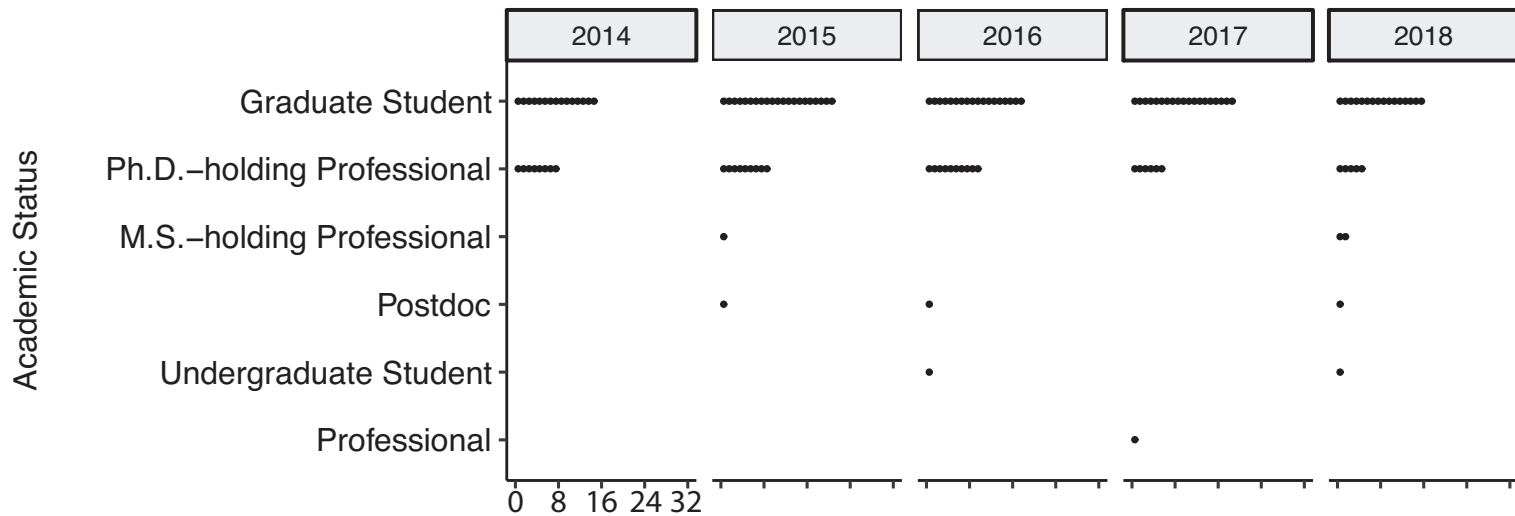
Attendee Age



Attendee Ethnicity



Attendee Academic Status



TIME	DAY 1	DAY 2	DAY 3	DAY 4	DAY 5	DAY 6	DAY 7
8:00 AM		Breakfast	Breakfast	Breakfast	Breakfast	Breakfast	
8:30							
9:00		Welcome lecture and assessment	BLAST				Invited Keynote
9:30							
10:00					Ecological Analysis	Sequencing Variants and Workflows	
10:30							
11:00		Computing Workflows	Fetching Data				Course Closeup and Assessment
11:30							
12:00 PM		Lunch	Lunch	Amplicon Sequencing	Lunch	Lunch	
12:30							
1:00							
1:30		Shell Programming	R Programming				
2:00					Statistical Analysis	Metagenome s	
2:30							
3:00		Moving Data	Data visualization				
3:30							
4:00							
4:30		Break and/or Office Hours	Break and/or Office Hours		Break and/or Office Hours	Break and/or Office Hours	
5:00				Break			
5:30							
6:00							
6:30		Dinner	Dinner	Dinner	Dinner	Dinner	
7:00							
7:30							
8:00	Meet and Greet	Invited Talk or Social Gathering	Invited Talk or Social Gathering	Invited Talk or Social Gathering	Invited Talk or Social Gathering	Invited Talk or Social Gathering	
8:30							
9:00							

I know how to process Illumina data

I understand what `per_library_stats.py` does

I know how to run R

I know the main differences in analyses offered by QIIME and mothur

I am familiar with `.biom` formatted files

I can name at least two different microbial metagenomic databases

I know what an R package is

I understand the structure of an OTU table

I know what a kmer is

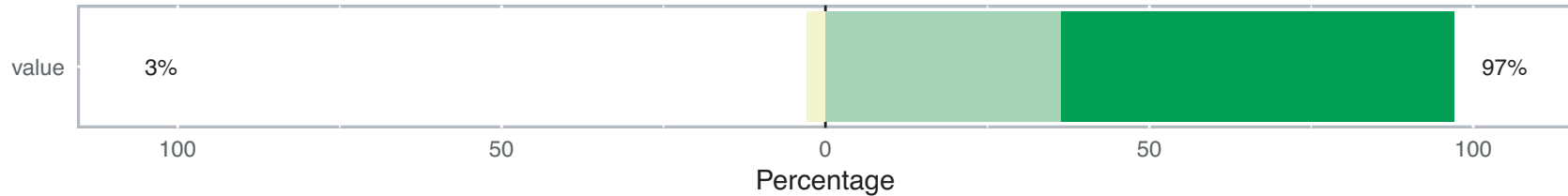
I know the difference between alpha and beta diversity

I know how to visualize microbial metagenomic data

I know how to use metadata to guide community analyses

I know how to assemble shotgun metagenomic data

Overall Rating Post EDAMAME Workshop, 2014 – 2018



Response



Very poor



Poor



Adequate



Good

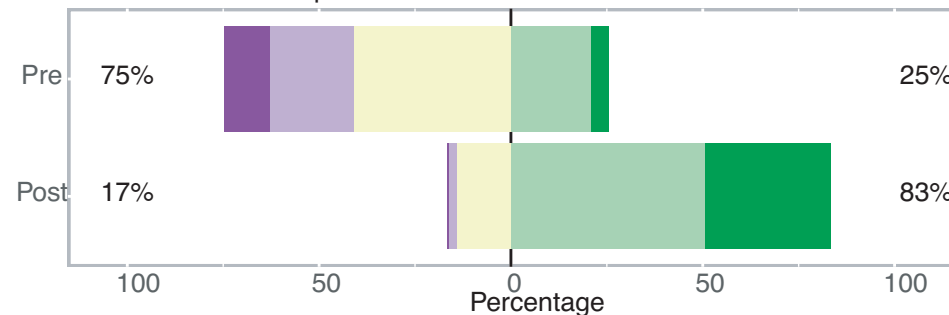


Very good

A. Comfort with Computational Tasks

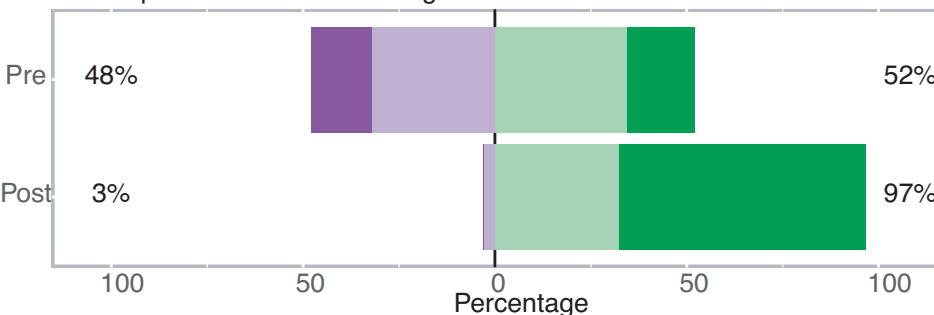
Response

- I do not know what this is.
- Very Uncomfortable
- Uncomfortable
- Comfortable
- Very Comfortable



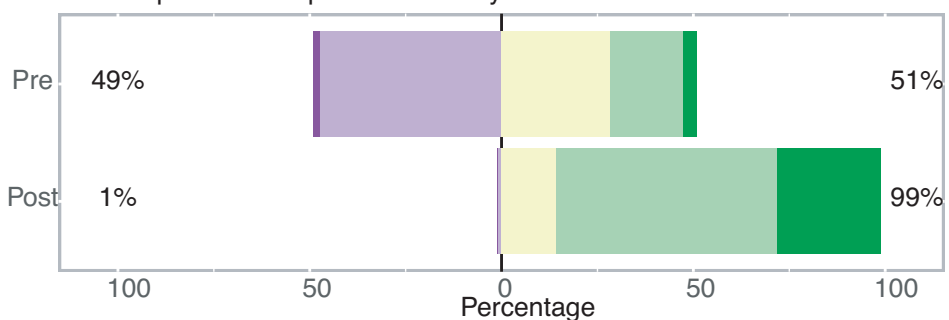
B. Computational Understanding

- Strongly Disagree
- Disagree
- Agree
- Strongly Agree



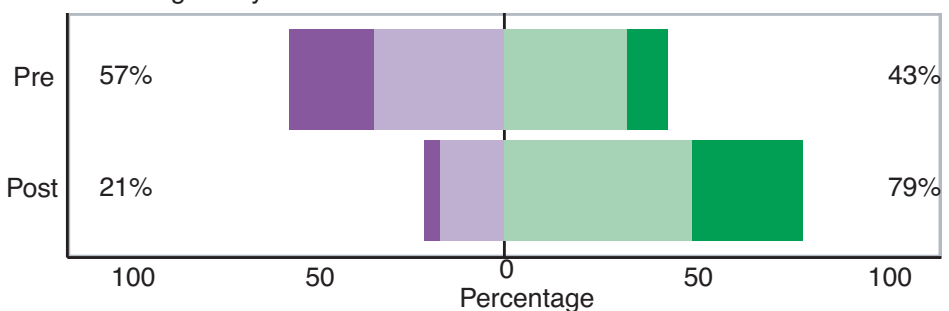
C. Perception of Computational Ability

- I'm not sure what this is
- No Ability
- Low Ability
- Intermediate Ability
- High Ability



D. Coding Ability

- Strongly Disagree
- Disagree
- Agree
- Strongly Agree



1. Regularly evaluate and change content to meet changing learner needs.
2. Maintain a high instructor to learner ratio.
3. Provide consistent workshop timing and fill the “middle-ground” duration needs of learners.
4. Understand the pros and cons of cloud computing for a workshop, and plan use of these resources well in advance.
5. Reach the broadest applicant pool of learners who have the potential to have the most gains from the training.
6. Consider the trade-off in workshop value (including instructor time) and maintaining economical costs to learners.
7. Plan well in advance to achieve best outcomes for applicants who require a US visa and international travel plans to attend the workshop.
8. Almost all learning engagement needs to happen on-site; efforts to engage learners pre-workshop were ineffective.
9. Scheduled classes should teach to the majority of learners to accomplish all our learning objectives. Office hours can help struggling learners catch up.
10. A welcoming and inclusive environment creates a positive workshop experience and is essential for effective learning.

Day 1	19:00-21:00
Day 2	8:00-9:00
	9:00-9:30
	9:30-10:30
	10:30-12:00
	12:00-1:00
	1:00-2:30
	2:30-3:00
	4:00-6:00
	6:00-7:30
	7:30-10:00
Day 3	8:00-9:00
	9:00-10:30
	10:45-12:00
	12:00-1:00
	1:00-2:00
	2:15-4:00
	4:00-5:00
	5:00-6:00
	6:00-7:30
	7:30-10:00
Day 4	8:00-9:00
	9:00-16:30
	6:00-7:30
	7:30-10:00
Day 5	8:00-9:00
	9:00-12:00
	12:00-1:00
	1:00-4:00
	4:00-6:00
	6:00-7:30
	7:30-10:00
Day 6	8:00-9:00
	9:00-10:00
	12:00-1:00
	1:00-4:00
	4:00-6:00
	6:00-7:30
	7:30-10:00
Day 7	8:00-10:30
	10:45-12:00

Table S1. An example of ε

Meet and Greet
Breakfast
Welcome lecture (code of conduct and learning objectives)
Assessment (front-end)
Best practices in computing workflows and using cloud computing
Lunch
Shell Programming
Moving data between local computers and the cloud
Break and/or Office Hours
Dinner
Invited Talk or Social Gathering
Breakfast
Comparing sequences with BLAST
Fetching data from federated databases (NCBI)
Lunch
Programming in R
Data Visualization in R
Afternoon Break
Optional lecture: students vote on topics of interest
Dinner
Invited Talk or Social Gathering
Breakfast
Amplicon sequence analysis (Mothur)
Dinner
Invited Talk or Social Gathering
Breakfast
Ecological analysis in R
Lunch
Statistical analysis in R
Break and/or Office Hours
Dinner
Invited Talk or Social Gathering
Breakfast
Sequence variant analyses and sequencing workflow pipelines
Lunch
Metagenomic analysis and assembly
Break and/or Office Hours
Dinner
Invited Talk or Social Gathering
Invited Talks (Keynote and Informal Panel)
Course close-up and Post-Assessment

an ideal one week schedule for EDAMAME. For a full schedule with links to 2018 EDAMAME tutorials and talks, s

's of the workshop are also available at <http://www.edamamecourse.org/materials/>.