INTRODUCTION

Motivated by a need to provide research opportunities to a growing number of STEM majors, we have developed a metagenomics research module, coupled with next generation sequencing (NGS), that allows students to address novel questions centered on urban microbial community diversity. Studies have shown that providing students with opportunities to investigate genuine research questions, as opposed to “cookbook” exercises, can result in improved learning outcomes and changes in students’ appreciation of science and research (1–8). We have successfully used the urban metagenomics module in an introductory microbiology lab at a large urban university to engage many more students in research projects than was previously possible using a traditional apprenticeship model. While we have focused on bacterial diversity and community dynamics in the urban environment, the module can be adapted for studying microbial communities in almost any environment and can be modified to target microorganisms other than bacteria as well as adapted to include whole genome shotgun sequencing and transcriptome analysis. The chance to participate in an innovative research project focusing on the urban environment resonates with our students because the urban environment is where they live, work, and study. We believe that using current techniques to investigate research questions that are relevant to our students leads to improvements in their attitudes toward science and improves their performance in the lab.

PROCEDURE

The metagenomics module is well suited for undergraduates working in small groups of three to six students. The protocol uses commercially available kits to reduce the preparation time for the lab and to maximize consistency. Working with the instructor, the students choose the site(s) to be investigated. The outline for the metagenomics module is as follows:

1. Identify sites and collect environmental samples by swabbing
2. Isolate and purify genomic DNA
3. Amplify 16S rRNA gene region using universal PCR primers
4. Analyze and quantify 16S rRNA amplicon
5. Roche/454 or Illumina sequencing of 16S rRNA amplicons
6. DNA sequence analysis

In this article, we cover the novel aspects of the protocol. A more detailed description of the protocol is provided in the supplemental materials. The detailed protocol, as well as a list of materials and reagents, are available at the CUNY Undergraduate Research Site (http://www.cuny.edu/research/sr/undergrad-research/for-faculty/AREM.html).

Sample collection

Collecting sufficient sample material from the urban environment can be a challenge because surfaces in the city often have fewer microorganisms than samples from soil, water, or humans, and the urban samples often contain contaminants that can inhibit PCR. Samples are collected using a swabbing technique. Students are instructed to swab an area roughly the size of their palm (~10 x 10 cm) and have successfully collected material from streets, sidewalks, subway platforms, playground equipment, and other common urban environments. Students swab three to six sites within a square meter and combine the material from the swabs into a single sample. Once the sample is collected on the swab, we use a modified version of the Mo Bio Power Soil kit protocol to extract the DNA from the sample.

Steps 2 to 5 are routine and described in detail in Appendix 1: Detailed protocol.

DNA sequence data analysis

The sequence data is run through filtering and analysis pipelines, such as QIIME (qiime.org), in order to define operational taxonomic units and is compared to bacterial 16S...
rRNA databases (such as Green Genes, http://greengenes.lbl.gov/cgi-bin/nph-index.cgi) to determine the bacterial diversity of the sample. Depending on the goals of the course and the experience of the instructor, the filtering and taxonomic assignments can be done with the students. Alternatively, these steps can be performed by the instructor outside of the class and the taxonomic classification and relative abundance data can be used in class. Many of the academic and commercial sequencing facilities will perform filtering and basic analysis of the raw sequence data. Classes that do not have a wet lab component can analyze data collected by students in other courses or data collected by other investigators.

Safety

The metagenomics module can be run safely in any lab that is designed to conduct routine molecular and microbiological techniques. At the sample collection step, a small fraction of the sample can be plated on media that support microbial growth. This allows for an excellent demonstration of the power of metagenomics because students see how little bacterial diversity they can identify by colony morphology on the plate compared to how many species they are able to identify using metagenomics. Because there is a risk of culturing pathogens collected in the environment, students should seal the plates with Parafilm after inoculating them and simply observe what they see growing on the sealed plate before autoclaving them.

CONCLUSION

We have used the metagenomics research module in the introductory microbiology lab sections over three semesters, reaching almost 400 students. Students have collected and analyzed data from a number of previously uncharacterized urban sites around New York City and we have begun to accumulate data over a number of time points that will allow students to determine how stable these bacterial communities are. Our preliminary outcomes assessment shows that the microbiology lab sections using the urban metagenomics module have lower rates of Ds, Fs, and withdraws relative to control sections (see Table 1). This may be the result of a greater engagement of students in the lab, due to the relevance and excitement of studying samples they have collected from within their community, which may carry over to keep students motivated to study and perform well throughout the course. In support of this, we have used post-course, anonymous, evaluation surveys to show that students in the lab sections using the urban metagenomics module tend to have more positive responses regarding the lab as a whole when compared to the responses from students in the traditionally formatted lab. For example, students in the sections using the metagenomics module report a greater confidence in interpreting scientific data and a greater appreciation for research (see Fig. 1).

Importantly, Figure 2 shows that students who participated in the urban metagenomics lab are more likely to remain involved in science, as measured by the number of students who continue on into masters, graduate, or professional programs in the biological and biomedical sciences. In summary, the urban metagenomics research module is an easily transferable, affordable unit that can be incorporated into a number of undergraduate STEM courses and used to improve student outcomes.

SUPPLEMENTAL MATERIALS

Appendix I: Detailed protocol

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REFERENCES

1. Buonaccorsi, V. P., et al. 2011. GCAT-SEEKquence: genome consortium for active teaching of undergraduates through increased faculty access to next-generation sequencing data. CBE Life Sci. Educ. 10:342–345.
2. Dolan, E. L. 2012. Next steps for vision and change: moving from setting the vision to change. CBE Life Sci. Educ. 11:201–202.
3. Handelsman, J., et al. 2004. Scientific teaching. Science 304:521–522.
4. Lopatto, D. 2007. Undergraduate research experiences support science career decisions and active learning. CBE Life Sci. Educ. 6:297–306.
5. Shaffer, C. D., et al. 2010 The genomics education partnership: successful integration of research into laboratory classes at a diverse group of undergraduate institutions. CBE Life Sci. Educ. 9:55–69.
6. Udovic, D., D. Morris, A. Dickman, J. Postlethwait, and P. Wetherwax. 2002. Workshop biology: demonstrating the effectiveness of active learning in an introductory biology course. Bioscience 52:272–281.
7. Wei, C. A., and T. Woodin. 2011. Undergraduate research experiences in biology: alternatives to the apprenticeship model. CBE Life Sci. Educ. 10:123–131.
8. Woodin, T., C. V. Carter, and L. Fletcher. 2010. Vision and change in biology undergraduate education, a call for action—initial responses. CBE Life Sci. Educ. 9:71–73.