The internal content review will include a roundtable to solicit input from external entities about the whole project.

The idea was suggested to include an infographic for case studies.

Working group members were asked to send any population-specific guidance they find.

Staff should collect current efforts by pharmaceutical companies.

Collect studies in which there are differences as a justification for why diversity in clinical trials matters, but the list doesn’t have to be exhaustive.

Megan also walked through the structure of the website that has been built in anticipation of hosting the content.

For population specific guidance, use well-documented cases of differences in pharmacogenomics based on ancestry to demonstrate why it’s important that these populations are included in biomedical research; the guidance is how to recruit specific populations to participate, not guidance on how patients with specific ancestry should be treated in a medical setting.
Best practices for community engagement include compensating participants for their time, effort, and expertise. However, compensation for community members is difficult through the state, as they would have to register as consultants.

**Next Steps:**
- Review case study content
- Develop content for population-specific guidance
- Continue building site
- Put together roundtable for external input