Title Interoperable, collaborative multi-platform variant calling with bebio

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Availability https://github.com/chapmanb/bcbio-nextgen

Documentation https://bcbio-nextgen.readthedocs.org/en/latest/contents/cwl.html

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bcbio (https://github.com/chapmanb/bcbio-nextgen) is an open, community effort to develop validated and scalable variant calling, RNA-seq and small RNA analyses. Last year at BOSC, we discussed our work to port bcbio's internal workflow representation to use the community developed Common Workflow Language (CWL: http://www.commonwl.org/). This transition removed barriers that prevented bcbio interoperability.

The practical benefit of changing to standardized workflow definitions is that bebio works on multiple heterogeneous platforms. Using CWL, bebio runs with Curoverse's Arvados (https://arvados.org/), UCSC's Toil (http://toil.ucsc-cgl.org/) and Seven Bridges' rabix bunny (http://rabix.io/). In addition, conversion to the Workflow Description Language (WDL: https://software.broadinstitute.org/wdl/) provides in-progress support for Broad's Cromwell (https://github.com/broadinstitute/cromwell) and DNAnexus's APIs (https://github.com/dnanexus-rnd/dxWDL). There is also ongoing work with other communities actively developing CWL integration, including Nextflow (https://github.com/nextflow-io/cwl2nxf) and Galaxy (https://github.com/common-workflow-language/galaxy).

Widespread bebio interoperability allows running in many computational environments without the overhead of maintaining bebio specific integrations. Users can run locally or on high performance computing clusters with schedulers like SLURM, SGE and PBSPro. In addition, CWL enabled runners work across the three major cloud providers: Amazon Web Services, Google Compute Engine and Microsoft Azure. Commercial platforms like Curoverse, Seven Bridges and DNAnexus enable clinical labs to run in controlled environments. The key component of this diverse support is collaboration through the CWL standard. This demonstrates the importance of community standard development, especially in research environments where it is typically difficult to fund maintenance of large scale infastructure development.

The talk will discuss the practicalities of adjusting below to use CWL and WDL. We balance infrastructure work for the transition to CWL with continued improvement of workflows and community support. Testing and documentation of below is more complex since we validate workflows, like germline and somatic variant calling, in many environments. This requires coordination between groups with different focus as platforms, analyses and standards develop. High level collaboration is increasingly important as we do more complex science, and we'll describe the role of the open bioinformatics community in enabling it.