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Title
Novel approaches in function-driven single-cell genomics

Permalink
https://escholarship.org/uc/item/013339ti

Journal
FEMS MICROBIOLOGY REVIEWS, 41(4)

ISSN
0168-6445

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Publication Date
2017-07-01

DOI
10.1093/femsre/fux009

Peer reviewed
REVIEW ARTICLE

Novel approaches in function-driven single-cell genomics

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One sentence summary: This review highlights the use of function-driven single-cell genomics for the targeted recovery of single-cell genomes from uncultivated or uncharacterized microbes implicated with a specific function or phenotype.

Editor: Antoine Danchin

ABSTRACT

Deeper sequencing and improved bioinformatics in conjunction with single-cell and metagenomic approaches continue to illuminate undercharacterized environmental microbial communities. This has propelled the ‘who is there, and what might they be doing’ paradigm to the uncultivated and has already radically changed the topology of the tree of life and provided key insights into the microbial contribution to biogeochemistry. While characterization of ‘who’ based on marker genes can describe a large fraction of the community, answering ‘what are they doing’ remains the elusive pinnacle for microbiology. Function-driven single-cell genomics provides a solution by using a function-based screen to subsample complex microbial communities in a targeted manner for the isolation and genome sequencing of single cells. This enables single-cell sequencing to be focused on cells with specific phenotypic or metabolic characteristics of interest. Recovered genomes are conclusively implicated for both encoding and exhibiting the feature of interest, improving downstream annotation and revealing activity levels within that environment. This emerging approach has already improved our understanding of microbial community functioning and facilitated the experimental analysis of uncharacterized gene product space. Here we provide a comprehensive review of strategies that have been applied for function-driven single-cell genomics and the future directions we envision.

Keywords: function-driven sequencing; single-cell genomics; microbial dark matter; single-cell activity

INTRODUCTION

Metagenomics has evolved through the need to better characterize our world by studying the composition and coding potential of complex microbial communities where a majority of the diversity is not yet cultivated. Facilitated by next-generation sequencing platforms, metagenomics can provide incredible sequence output from a minimally disturbed complex environmental sample, making it the gold standard for recovering DNA sequences from samples of interest. Despite the magnitude of the output, the logistics of metagenome assembly are complicated and many community features can be lost within the sequence data. Furthermore, annotating assembled sequences depends on homology-based computational tools to infer function with varying degrees of ‘homology creep’ (Woyke and Jarett 2015), and provides no information regarding the activity of that organism within the environment unless coupled with experimental approaches like metatranscriptomics, proteomics or stable isotope probing. Even when an ‘all of the above’ approach is taken, the final result is the average of a bulk population, masking the large variability in functional activity due to spatial microenvironments or phenotypic noise known to drive divergent functions even in clonal populations (Ackermann 2013). Thus, the strength of metagenomics for its indiscriminate capture of all DNA from a heterogeneous sample is also its pitfall in that the complex output can confound the recovery of...
complete genomes, obscure genomic population heterogeneities (Engel, Stepanauskas and Moran 2014), overlook rare organisms (Ainsworth et al. 2015), miss ecological relationships, blend spatially divergent populations and diminish perspective for the interpretation of recovered sequences of unknown function.

Single-cell genomics offers a complimentary approach to metagenomics by capturing and amplifying DNA from a single isolated cell, as opposed to many cells in metagenomics, and is routinely capable of producing partial (Zhang et al. 2006), and even complete (Woyke et al. 2010), non-composite genomes. This approach thus addresses some of the caveats of metagenomics at the sacrifice of throughput. Although technical limitations and inherent biases currently exist for single-cell approaches (Lasken 2012; Gawad, Koh and Quake 2016), within microbiology single-cell genomics has demonstrated itself as a powerful tool by generating genomes from rare (Martijn et al. 2015), symbiotic (Sieg et al. 2011) and previously uncharacterized microbial lineages (Marcy et al. 2007; Blainey et al. 2011; Rinke et al. 2013). In addition, auxiliary DNA from viruses, organelles, plasmids, transformed DNA and symbionts is also captured and sequenced along with the host DNA due to its colocalization with the cell (Stepanauskas 2015). Thus, associations between these genetic elements and the host organism of interest are maintained with high resolution, allowing characterization of subtle ecological interactions (Yoon et al. 2011; Roux et al. 2014). When coupled with microscopy, single-cell genomics enables spatiotemporal characterization of microscale environments, improving the resolution at which our understanding of microbial ecology takes place (Landry et al. 2013). Single-cell genomics has identified a novel genetic code (Campbell et al. 2013), uncovered unexplored protein sequence space relevant for biotechnology and human medicine (Wilson et al. 2014), and facilitated the expansion of novel branches in the bacterial and archaeal tree of life and improved phylogenetic read anchoring for metagenomic data sets (Rinke et al. 2013). With these efforts, pipelines for the single-cell isolation and sequencing of environmental microbes have optimized their efficiency and throughput (Hutchison et al. 2005; Woyke et al. 2011; Rinke et al. 2014), thereby advancing single-cell genomics methods to the next level.

The major outstanding limitation in metagenomic and single-cell approaches is the overreliance on using predicted protein function as a proxy for functional activity in the environment. These approaches lack the ability to discern which recovered populations are active, potentially overestimating the importance of abundant organisms while overlooking significant ecological contributions of lower abundance organisms (Martinez-Garcia et al. 2012). This caveat becomes even more pronounced when trying to study ‘microbial dark matter’ i.e., microbes that lack characterized cultured representatives (Rinke et al. 2013; Kamke et al. 2014; Brown et al. 2015) and whose genes often lack functional prediction (Youssef et al. 2011; Kantor et al. 2013; McLean et al. 2013). Thus, although both metagenomic and single-cell genomic approaches can recover novel genes, neither approach currently provides insight into the function of the gene or activity level of the organism.

Identifying the function of a gene has traditionally relied on classical genetic knockout/complementation studies, and more recently high-throughput relative fitness studies (Wetmore et al. 2015). Because these approaches require compatible genetic toolkits and cultivable organisms, they are unsuitable for studying the function of unknown genes from the uncultivated majority. Function-based screens of metagenomic DNA sequences have previously utilized clone libraries to heterologously express environment-derived DNA fragments in a gain-of-function approach (Daniel 2005). Escherichia coli, the workhorse of heterologous screening approaches, was computationally determined to be able to transcribe ~40% of genes from well-known cultivated lineages of microbes with the typical expression library approach (Gabor, Alkema and Janssen 2004). Considering downstream incompatibilities such as codon bias (Tuller et al. 2010), strategic rare codon utilization (Komar et al. 1999), required metabolite pools, post-translational modification, accessory proteins, apoprotein activation, secretion and even the availability of a compatible read-out assay, it becomes apparent that only a small fraction of functional space is accessible through this approach. Furthermore, when introducing more divergent DNA sequences, such as those from candidate phyla, the success rate for accessing the vast functional diversity that exists with current tools is expected to rapidly diminish. Thus, the majority of unknown functions from uncultivated organisms remain obscured within their native expression hosts.

Due to the increasing recovery and accumulation of sequences of unknown function, a context-driven approach to investigate the roles of these genes within their native hosts and environment is required. This will facilitate improved protein annotation, interpretation of microbial networks and understanding of microbial influences on biogeochemistry (Hicks and Prather 2014; Woyke and Jarett 2015). Thus, an increased focus on characterizing the functional roles and activity levels of uncultivated organisms in conjunction with downstream genomic sequencing has motivated the development of methods to ascertain both functional trait and/or phenotype, and corresponding coding potential from these organisms at the resolution of a single cell. Here we present the approaches and limitations of pioneering studies focused on developing a suite of methods for the function-driven single-cell identification, isolation and sequencing of uncultivated environmental microbes. Though this review focuses on these approaches from a largely bacterial and archaeal perspective, many of the same strategies can be adopted to small environmental eukaryotic cells, and some are being applied in human cells to address human health issues (Gao et al. 2004; Yu et al. 2011).

FUNCTION-DRIVEN APPROACHES

Label-free

The earliest ‘targeted’ single-cell screens took advantage of previously characterized environments where distinguishing morphological traits were used as identifying selection criteria. Marcy et al. first applied this approach in 2007 for single-cell genomics from members of the uncultivated TM7 phylum (Marcy et al. 2007). To isolate their target, rod-shaped cells from the human mouth were sorted and enriched in a microfluidic chip (Rodrigue et al. 2010). Other morphological features relating directly to function, including photosynthetic pigments (Rodrique et al. 2009), polyhydroxybutyrate production (Tyo, Zhou and Stephanopoulos 2006) and magnetotaxis (Kolinko et al. 2012), have also been used as selection criteria for identifying and isolating single cells in true function-driven screening approaches (Fig. 1). However, since a majority of activities of interest cannot be selected by a unique morphological trait,
Label-based screens provide an alternate route to visualize additional functions.

**Label-based approaches**

Recent innovations in function-driven single-cell genomics have successfully exploited the association, metabolism or integration of labeled substrates to implicate functional activity in specific microbial guilds within environmental communities. These studies have utilized fluorescently tagged (Martinez-Garcia et al. 2012; Geva-Zatorsky et al. 2015) and isotopically labeled (Huang et al. 2009; Li et al. 2012) substrates (Fig. 1) to identify and recover individual active microbes that respond to, and interact with, these substrates.

**Fluorescent substrate**

The fluorescent substrate single amplified genome analysis (FS-SAGA) method was developed by Martinez-Garcia et al. (2012) to identify populations active in the degradation of the complex carbohydrate laminarin in aquatic systems. Though bacterial degradation of this abundant substrate had been demonstrated in aquatic environments (Arnosti 2011), the identity of the responsible population remained obscure (Alderkamp, van Rijssel and Bolhuis 2007). To conclusively identify this population, Martinez-Garcia et al. spiked fluorescently labeled laminarin into freshly recovered aquatic samples. The samples were then screened by flow cytometry to recover cells that became increasingly fluorescent (strategy outlined in Fig. 1). The captured cell population was largely dominated by a few Verrucomicrobia taxa, which had low overall relative abundances (<1%) within the starting population as determined by culture-independent methods. Single-cell genomes generated from this population were highly enriched in a number of glycoside hydrolase families, suggesting the accuracy of the FS-SAGA method for identifying this highly active, yet relatively rare, population and implicating the activity of these specific hydrolases with laminarin degradation.

**Bioorthogonal tagging**

A caveat of fluorescently labeled substrates is that the modification required to attach fluorophores may abolish native recognition, transport and metabolism of that substrate. One strategy to circumvent the complication of bulky fluorophore attachment has been the utilization of non-canonical substrates that instead contain only a small bioorthogonal tag. A bioorthogonal reaction, such as the commonly used azide-alkyne ‘click’ reaction, is advantageous since it readily proceeds within biological systems without cross-reaction (Baskin et al. 2007). During incubation, these non-canonical substrates are integrated into the biomass of metabolically active, but potentially non-replicative, cells allowing incorporation into cellular structures before a subsequent introduction of a ‘click’ compatible fluorescent marker (Speers, Adam and Cravatt 2003) (Fig. 2, no. 7). This has been successfully demonstrated with non-canonical amino acids, nucleotides, lipids and sugar analogs modified to contain either an azide or alkyne group for ‘click’ chemistry compatibility (Dieterich et al. 2007; Salic and Mitchison 2008; Neef and Schultz 2009; Paredes and Das 2011; Hatzenpichler et al. 2014; Samo et al. 2014; Geva-Zatorsky et al. 2015), though the enzymatic
promiscuity that permits integration of these analogues may not be universal among all bacteria. Hatzenpichler et al. (2016) applied a bioorthogonal non-canonical amino acid tagging (BONCAT) approach to investigate the activity and microbial ecology of slow-growing methane-oxidizing/sulfur-reducing archaea/bacteria consortia with intra-aggregate resolution. Methane seep enrichment cultures were incubated, with and without methane stimulation, in the presence of L-homopropargylglycine (HPG). This alkyne-containing non-canonical amino acid is a structural analog for L-methionine that is readily taken up and integrated into nascent proteins within active cells. Following incorporation of HPG into cellular proteins, click-based fluorophore labeling marked ~25% of aggregates as translationally active (compared to ~2% in samples incubated without methane). The community identity of each individual aggregate was then determined by sorting the fluorescent aggregates with flow cytometry, followed by lysis, genome amplification with Phi29 polymerase (as detailed in Fig. 1), and amplification of the 16S rRNA gene with ‘universal’ primers. This approach allowed mapping the taxonomic identities of six distinct archaeal lineages (ANME-1a, 1b, 2a, 2b, 2c and 3) in individual aggregates with their preferred syntrophic sulfate-reducing partners.

**Substrate-independent isotope**

To avoid complications of modified substrates entirely, isotopically labeled substrates can be introduced instead. However, because of the cost and logistical challenge associated with synthesizing substrates from stable isotopes, Berry et al. (2015) developed a substrate-independent strategy using deuterated water for activity-based identification of individual cells following perturbation of a community with a native substrate. As microbes grow in the presence of heavy water (D2O), deuterium is actively integrated within their biomass through the synthesis of new lipids and proteins. The replacement of the C-H bond with the C-D bond causes a spectral shift due to the change in vibrational energy when measured by Raman microspectroscopy (Wei et al. 2013). Thus, the emergence of the C-D peak in the Raman spectrum serves as a proxy for the extent of biosynthetic activity, thereby unambiguously labeling all active organisms in a substrate-independent manner (Kopf et al. 2015). This strategy enables a targeted approach when obtaining labeled substrates of interest is prohibitive, or where environmentally relevant substrates are unknown. Berry et al. (2015) coupled the labeling of both pure culture and complex environmental microbial cultures with D2O and employed single-cell Raman microspectroscopy detection within capillary tubes (Fig. 1) to screen and sort individual cells. Following benchmarking of the method with cultures of model organisms from diverse phyla, the approach was applied to a mouse cecal microbiome stimulated with a range of carbohydrates. Responses from randomly screened cells following incubation ranged from 9% (no substrate addition) to 41% (mucin) and 81% (glucose) labeling, suggesting that different substrates successfully activated different subpopulations. To validate the observed result, fluorescent in situ hybridization (FISH) probes targeting two known mucin degraders served as positive controls for activity and correctly identified their targets as active under expected conditions. This study demonstrates the power of the substrate-independent isotope approach by making nearly any substrate and environment amenable to probing.

**Activity-based probes**

Though largely yet to be applied, there are many function-based probes compatible with application in single-cell genomics that originate from, and elaborate on, the field of activity-based profiling (ABP). ABP historically focused on the identification of a specific protein family activity from within a bulk protein pool isolated from a single organism (Barglow and Cravatt 2007). The most extensive developments in ABP have been employing family-specific probes for the identification of differential protein activity levels between healthy and diseased states (Jessani and Cravatt 2004). These probes have chiefly targeted large protein families including serine (Liu, Patricelli and Cravatt 1999), cysteine (Kato et al. 2005) and metallohydrodases (Saghatelian et al. 2004), and have resulted in numerous insights in proteome function and annotation (Adam et al. 2004). Strategies for improving the design and activity of these probes have been extensively reviewed (Cravatt, Wright and Kozarich 2008) and include methods for improving reactivity (Walvoort et al. 2012), quantification (Okerberg et al. 2005), labeled fraction recovery (Chan et al. 2004), identification of participating proteins and active site residues (Speers and Cravatt 2005), and effectiveness of target competitive inhibitors (Evans et al. 2005).

Activity-based probes are typically designed using a modular three-component strategy: a warhead for target specificity, a tag for detection, and a polyfunctional linker for joining the warhead and tag that can also aid in quenching and/or covalent attachment following activation (Sadler and Wright 2015) (Fig. 3). Two primary classes of functional probes, mechanism and affinity based, have been developed and successfully demonstrated (Chauvigne-Hines et al. 2012). Mechanism-based probes rely on the direct enzymatic transformation of the probe by a specific cellular enzyme or metabolite to become activated. These probes often mark their target with high specificity by forming a covalent bond with a key active residue in the enzyme, functioning as an irreversible inhibitor (Walvoort et al. 2012). This approach is advantageous when looking to characterize only active protein forms from zymogens (Adam et al. 2004). Affinity probes alternatively do not require mechanistic activation by the target, but physically associate with a receptor or molecule of interest for retention or tag activation (Chan et al. 2004). An example of this strategy is the nanoparticle-based quorum-sensing...
probe that relies only on the affinity between the homoserine lactone warhead and the intracellular quorum-sensing receptor (Mukherji et al. 2013) (Table 1). In this example, however, no strategy for tag activation or covalent attachment exists upon receptor binding, likely reducing the sensitivity of the probe screen when advancing to environmental samples.

Similar to how probing enzyme targets from a bulk pool is a powerful approach as it can greatly reduce the complexity of results when screening the proteome, activity-based probes can analogously be applied to a complex microbial community to return a narrowed result of what microorganisms in the given environment are actively performing an activity of interest (Fig. 2). Translating this technology from its previous proteomic application to single-cell thus shifts the focus from a family of proteins within a bulk pool as the fundamental unit of activity to the functional profile of an individual cell within a community as a fundamental unit. Though applications in activity-based profiling of environmental microbes are just emerging (Sadler and Wright 2015), we envision that by coupling this approach with cell sorting and single-cell genomics, activity-based probe selection offers a strategy to parse a complex community, identify individual organisms participating in a given function and relate this activity, with the context of their genome, to the community at large. In addition to surveying individual cellular activities with a specific functional probe, screens can be multiplexed with other cellular function or status signals using a suite of probes developed for cellular conditions and activity levels (Sieracki, Cucci and Nicinski 1999; Kalyuzhnaya, Lidstrom and Chistoserdova 2008; Chan, Dodani and Chang 2012) (Table 1).

**Magnetic capture**

Beyond magnetic enrichment of magnetotactic bacteria as a function-driven screen (Kolinko et al. 2016), magnetic capture using functionalized magnetic beads presents itself as a high-throughput function-driven selection screen with the ability to enrich all members with a given extracellular marker from an entire sample (Fig. 2, no. 2). The ability to recover cells using aptamer (Chang et al. 2013) or phage binding domains (Kretzer et al. 2007) has already demonstrated that this approach could readily be applied for function-driven single-cell genomics.

**CHALLENGES**

Beyond increasing the array of compatible functional approaches, one of the primary obstacles to improving the application of function-driven single-cell screens within environmental samples is ensuring a sufficiently strong probe signal beyond that of background noise e.g. natural phycobiliproteins can produce higher fluorescence than organic fluorophores (Chiu 2014). Due to the limited number of targets for tagging within a cell (e.g. receptors or enzyme active sites for a specific function of interest), ensuring conclusive detection of a positive result beyond background noise, whether fluorescent or Raman, is paramount to the effectiveness of the single-cell approach. Though general strategies for improving probe design have been widely studied and extensively reviewed (Chan, Dodani and Chang 2012; Vendrell et al. 2012), coupling the addition of these probe improvements with the specificity required for function-driven screens remains largely a case-by-case approach. Here we outline some of the general strategies that have been successfully used to improve the detection of function-driven activities.

**Activity-based probes**

**Turn on signal**

Minimizing avoidable background signals from non-specific associations of fluorescent probes have resulted in the design of
### Table 1. Potential strategies for single-cell identification and targets of interest. Numbering corresponds to labeling targets in Figure 2.

<table>
<thead>
<tr>
<th>Target</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Labeling strategies</strong></td>
<td></td>
</tr>
<tr>
<td>(1) Extracellular integration</td>
<td>Oligosaccharide labeling (Geva-Zatorsky et al. 2015)</td>
</tr>
<tr>
<td>(2) Extracellular recognition</td>
<td>DNA aptamer (Chang et al. 2013), phage binding domain (Kretzer et al. 2007)</td>
</tr>
<tr>
<td>(3) Extracellular enzyme affinity</td>
<td>Fluorescent substrate (Martinez-Garcia et al. 2012), glycose hydrolases (Chauvigne-Hines et al. 2012), phytase (Berry and Harich 2013), penicillin binding proteins (Kocaoglu and Carlson 2013), β-lactamases (Bottcher and Sieber 2012; Shao and Xing 2012), showdomycin (Bottcher and Sieber 2010)</td>
</tr>
<tr>
<td>(4) Intracellular receptor</td>
<td>Quorum sensing (Mukherji et al. 2013)</td>
</tr>
<tr>
<td>(5) Intracellular product</td>
<td>Polyhydroxybutyrate granules (Yio, Zhou and Stephanopoulos 2006), hypochlorous acid production (Sun et al. 2014), ROS production (Gomes, Fernandes and Lima 2005)</td>
</tr>
<tr>
<td>(6) Intracellular enzymatic reaction</td>
<td>Non-ribosomal peptide synthetases (Kono et al. 2015), polyketide synthases (Meier et al. 2009), nucleophilic RNAs (McDonald et al. 2014), adenylation (Duckworth et al. 2012), glyceraldehyde-3-phosphate dehydrogenase (Kaschani et al. 2012), glutathione reductase (Lou et al. 2014), sulfatase (Park, Rhee and Hong 2012), phosphatase (Kim et al. 2011), aldehyde dehydrogenase (Adam, Cravatt and Sorensen 2001), palmitoyl acetyltransferases (Zheng et al. 2013), β-glucosidases (Kallemeijn et al. 2012; Walvoort et al. 2012), agmatine deiminase (Marchenko et al. 2015), monoamine oxidase (Li et al. 2014), N-acetylgalactosaminidase (Kalidasan et al. 2013), alkaline phosphatase (Kim et al. 2011), metalloproteases (Sieber et al. 2006), cysteine proteases (Kato et al. 2005), serine hydrolase (Liu, Patricelli and Cravatt 1999)</td>
</tr>
<tr>
<td>(7) Intracellular integration</td>
<td>ACNOMAT (Dietrich et al. 2007), degenerate tRNA synthetases (Ngo et al. 2009), RNA labeling (Paredes and Das 2011)</td>
</tr>
<tr>
<td>(8) Isotope integration</td>
<td>2H2O (Berry et al. 2015), 13CO2 (Li et al. 2012), [13C] naphthalene (Huang et al. 2009)</td>
</tr>
<tr>
<td><strong>Characteristics</strong></td>
<td></td>
</tr>
<tr>
<td>General properties</td>
<td>Magnetotaxis (Kolinko et al. 2012), cell morphology (Marcy et al. 2007)</td>
</tr>
<tr>
<td>Intracellular ions</td>
<td>pH (Han and Burgess 2010), [Zn2+] (Wallkup et al. 2000)</td>
</tr>
</tbody>
</table>

‘turn-on’ probes that display improved signal intensity only following structural or conformational activation (Fig. 3). For fluorescent probes, this can be achieved with quenching modules that absorb the fluorescent reporter signal (Li et al. 2014), fluorogenic precursor probes (Weissleder et al. 1999; Kim et al. 2011), photoinduced electron transfer (de Silva, Moody and Wright 2009), structural conformation (Mello and Finney 2001) and a variety of other strategies (Chan, Dodani and Chang 2012). In each case, a covalent or conformational modification to the probe through a specific activity of the cell results in an increase in detectable signal, localizing the activated probe fluorescence to within functionally active cells.

**Turnover vs non-turnover**

Two primary types of mechanism-based probes have been described with regard to activation: turnover and non-turnover. Non-turnover probes are essentially irreversible inhibitors that occupy the enzymatic active site following activation. These probes often function through high-affinity covalent bonding to active site residues that result in a structural change for probe-signal activation (Kallemeijn et al. 2012). Although this probe strategy ensures the activated fluorophore is retained inside the cell, this approach imposes a maximum signal for each cell due to the finite number of targets. In addition, regardless of enzyme kcat or km (the parameters that would largely determine relative cellular activity levels), if the concentration of target enzyme is present at low levels within the cell, detecting a discernable signal above background could be prevented. This limitation thereby prevents a relative comparison in true activity levels between different organisms as measured by the fluorescent signal in the function-driven screen. Turnover probes, alternatively, do not result in inactivation of the target enzyme following modification and thereby allow repeated activation of multiple probe molecules by a single enzyme (Kalidasan et al. 2013). This strategy improves the maximum theoretical signal attainable by a given cell by maintaining a functional pool of enzymatic activity independent of probe activation. In addition, kcat and km could influence the rate and concentration to which the activated probe can accumulate, allowing semiquantitative insights into relative activity levels between cells.

**Retaining signal**

Permeable probes are required to pass through intact membranes to reach intracellular targets of interest in physiologically active cells. Following activation, however, turnover mechanism-based and affinity probes often remain free to diffuse out of the cell into the bulk environment, diminishing the intensity of the signal within the cell and increasing background. While non-turnover probes are retained in the cell through their mechanistic bonding to the enzyme active site, if activated turnover probes could be retained inside the cell through non-specific covalent bonding to cell biomass instead of the enzymatic site, this approach would allow for both the covalent retention of the probe within a cell of interest while allowing the target enzyme to continue activating probes (Kalidasan et al. 2013). Alternatively, activated probes that become unable to diffuse through the membrane due to changes in charge or solubility are other strategies that could be designed for retaining signal. Using classical probe design approaches, those structures that can integrate modular components to address all the above design strategies including turn-on signal, enzyme active site turnover, and a strategy for intracellular probe retention present a route for best chances for detection of signal above noise in a function-driven detection screen (Kalidasan et al. 2013).
Other challenges

Modern fluorophores
Many of the fluorescent markers conjugated to functional probes have remained classic organic dyes (e.g. fluorescein, rhodamine) although advances in materials science have produced new fluorescent molecules with improved properties. Examples include semiconductor quantum dots (Qdots) (Gao et al. 2004) and x -conjugated polymer dots (Pdots) (Wu et al. 2008). This limitation for function-driven single-cell genomics from classical dyes stems from their relatively low absorbivity and poor photostability (Chiu 2014). Recent advances in dot fluorophore technology have improved the absorbivity and quantum efficiency of applied probes and have even demonstrated compatibility with click chemistry labeling (Wu et al. 2010).

Surface-enhanced Raman detection
Raman microspectroscopy is a very powerful tool as it provides an entire cell ‘fingerprint’ by measuring inelastic scattering of photons from abundant bonds within a single cell. However, Raman vibrations are relatively rare events in cells, with only 1 in 10^6–8 photons causing a detectable vibration, and often require measurements on the order of minutes per cell (Jarvis and Goodacre 2008). In addition, due to the complexity of simultaneously probing all vibrational energies within the cell, specific signals from within the spectrum can be convoluted. The addition of nanometer-scale Ag/Au particles or colloids to a culture of interest results in a surface-enhanced Raman spectroscopy (SERS) phenomenon, whereby the sensitivity and acquisition time are improved in isotopic measurements on single cells (Kubryk et al. 2015). SERS not only enables improved throughput of the approach, but also potentially reduces the amount of isotope incorporation necessary for detection, minimizing any negative impacts of high levels of isotopically labeled substrates. Current limitations with the SERS approach within function-driven single-cell screens include heterogeneity of nanoparticle binding (Kubryk et al. 2015), and any potential toxicity due to nanoparticle association with cells (Rai, Yadav and Gade 2009).

Sample activity
Unlike traditional metagenomics or single-cell sampling methods where samples can be taken and immediately preserved through freezing or fixation as instantaneous snapshots, function-driven methods require both viability and activity to be maintained following sampling. Because of this, and in the interest of maintaining an as close to in situ observation as possible, minimal disturbance to the physical and chemical properties of the sample is necessary. In addition, delays in sample screening perturb the natural abundances and activities of the microbial community within the sample, displacing events from the true in situ activity. These limitations logistically restrict the extent of samples that can be screened to those that can be quickly recovered with minimal disturbance from the target environment to the lab.

Disaggregation of cells
As the name implies, single-cell genomics requires the ability to manipulate individual cells, a task which is not always trivial depending on sample type. Vortexing, sonication, aspiration, centrifugation and even grinding are all approaches that have been used to liberate single cells from difficult sample environments (Rinke et al. 2014). The optimal method for liberating cells while still maintaining activity necessary for downstream screening will vary by sample type and likely require a case-by-case approach.

Sample fixation
Fixation is often a critical step for the preparation of samples where membrane permeabilization is required for introduction of a probe or label. For function-driven single-cell genomics this process can become problematic, as fixation often interferes with downstream whole genome amplification. Formaldehyde is a common fixation agent for the stabilization and inactivation of cells through global crosslinking. However, crosslinking of genomic DNA with formaldehyde has been shown to prevent amplification (Ben-Ezra et al. 1991), alter nucleotide sequences and degrade DNA (Williams et al. 1999), making it incompatible with single-cell genomics. While ethanol, a precipitating fixative, has been found to produce fixed cells that yield sufficient amplification product for single-cell sequencing, it comes at the cost of reduced genome recovery (Clingenpeel et al. 2014). A range of alternative fixation agents and protocols with varying degrees of previously reported DNA amplification success could be screened with Phi29 for compatibility with whole genome amplification (Srinivasan, Sedmak and Jewell 2002).

Cell lysis
Lysing an isolated cell under mild enough conditions to maintain the integrity of the genomic DNA prior to whole genome amplification can be challenging due to the varying degree of difficulty lysing structurally diverse cells. Alkaline lysis, lysozyme treatment, heat, freeze-thaw and detergents have been used with varying success of subsequent amplification by method and cell type (Rinke et al. 2014). Developing efficient cell lysis methods that routinely liberate DNA from a wide range of microbial cells will broaden potential targets of single-cell approaches.

FUTURE
While many compatible tools and methodological strategies outlined here for probing the function of the uncultivated majority exist, only a handful of remarkable studies to date have applied these tools for the characterization and recovery of single-cell genomes from the environment. These findings, such as the high levels of functional activity from low abundance keystone organisms and the resolution of taxonomic partnerships within diverse methane-oxidizing aggregates, have contributed to the fundamental understanding of microscale ecological dynamics. Continuing to apply existing tools and developing new approaches for characterizing the in situ function of uncultivated microbes in the environment will continue to expand our understanding of the functional role of ‘microbial dark matter’. Moving forward, one primary pitfall that remains with this strategy is that linking the activity with a particular coding sequence can be easy to apply if the encoded function is homologous to known and previously characterized genes. If recovered genes involved in the targeted function screen are so divergent that they can only be assigned as hypothetical, attributing the screened activity to specific sequences in the genome becomes obfuscated. However, compiling recovered genes of unknown function implicated with the screened activity and coupling them with an optimized DNA synthesis and expression approach could allow the targeted discovery of truly novel gene function from uncultivated organisms.
SUMMARY

Function-driven single-cell genomic approaches offer a unique route to directly study the activities of uncultivated organisms, at single-cell resolution, in an in situ style approach without any prior knowledge of cellular activity. The ability to screen all members of a microbial community, independent of our ability to cultivate them, and implicate individual organisms as actively involved with a specific functional activity or exhibiting a certain morphological phenotype provides valuable information regarding their ecological role within a given ecosystem. Being able to then couple the genomic sequence of the identified organisms back to the screened activity or trait allows for the characterization of potentially unknown and unstudied genes and pathways in their native hosts. This represents a critical strategy to advance our understanding of microbial community functioning and move beyond purely sequence-based predictions.

FUNDING

The work conducted by the U.S. Department of Energy Joint Genome Institute, a DOE Office of Science User Facility, is supported under Contract No. DE-AC02-05CH11231. This work was supported under the LBNL Microbes to Biomes LDRD entitled “function-driven” genomics’.

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ACKNOWLEDGMENTS

The work conducted by the U.S. Department of Energy Joint Genome Institute, a DOE Office of Science User Facility, is supported under Contract No. DE-AC02-05CH11231. This work was supported under the LBNL Microbes to Biomes LDRD entitled “function-driven” genomics’.

Conflict of interest. None declared

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