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Metabolomics tools for the synthetic biology of natural products

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Abstract

Metabolomics plays an increasingly central role in within the Design – Build – Test cycle of synthetic biology, in particular in applications targeting the discovery, diversification and optimized production of a wide range of natural products. For example, improved methods for the online monitoring of chemical reactions accelerate data generation to be compatible with the rapid iterations and increasing library sizes of automated synthetic biology pipelines. Combinations of label-free metabolic profiling and 13C-based flux analysis lead to increased resolution in the identification of metabolic bottlenecks affecting product yield in engineered microbes. And molecular networking strategies drastically increase our ability to identify and characterize novel chemically complex biomolecules of interest in a diverse range of samples.

Graphical abstract

Synthetic biology of natural products

Synthetic biology facilitates the biosynthesis of pharmaceutical ingredients and other high-value chemicals by employing the Design – Build – Test cycle of engineering to guide the systematic enhancement of microbial factories [1-4]. Exemplary successful applications of synthetic biology to natural product production include a one-pot method for menthol biosynthesis in *Escherichia coli* [5], the modular extension of a styrene biosynthesis pathway to produce 2-phenylethanol [6], cannabinoid biosynthesis in yeasts [7], and the heterologous production of antibiotics using extensively refactored biosynthetic gene clusters: myxobacterial α-pyrone antibiotics in *Myxococcus xanthus* [8] and kasugamycin (an aminoglycoside antibiotic isolated from *Streptomyces kasugaensis*) in actinomycetes [9]).

A recent review by Smanski and colleagues [10] provides details of recent advances in the technologies underpinning the Build aspects of the synthetic biology cycle, including pathway construction and pathway screening, while a complementary review by Chen *et al.* [11] focuses
on the modelling approaches for the construction and optimisation of cell factories for bio-
production, which cover a large part of the Design activities. In the present review, we will in
turn focus on the Test component of synthetic biology, focusing in particular on advances in
metabolomics as a discovery and debugging tool for metabolically enhanced microbial systems.

Test Analytics – Appropriate Technologies

Mass spectrometry (MS) coupled to chromatography remains the domineering technology used
for the quantification of natural product targets and is also the most widely used platform for
the global profiling of the impact of an engineered biosynthetic pathway on the microbial
metabolome. The challenge for the analytical technologies is to achieve the acquisition speed
and sensitivity required to meet the high-throughput needs of a synthetic biology-based
pipeline. Traditionally, products are measured directly from an aliquot of cell culture medium
or – in the case of volatile products – they are captured in solvent overlays and transferred to
vials or multi-well plates for analysis. These approaches are often slow (tens of minutes per
sample plus preparation time) and provide only a snapshot of what is occurring at a given time.

To overcome this limitation, much effort has been invested into the development of improved
methods for the online monitoring of chemical reactions; which would provide greater control
of sampling and provide dynamic results with regards to product turnover. Definitions of the
analytical terminology described herein are summarised in Table 1. In recent work by Yan et al.,
desorption electrospray ionization (DESI) coupled to ion-mobility mass spectrometry was used
for the high-throughput screening of biocatalysis directly from bacterial colonies on agar plates
[12], which can in principle be applied to a broad range of substrates and products, including
free amines, carboxylic acids, alkaloids and phenols; multiple analytes can be detected in a
single analysis thus allowing for the screening of diverse strain libraries with complex product
profiles. DESI-MS was also applied to the rapid analysis of enzyme kinetics by Cheng and co-
workers [13], who measured product formation in a buffered aqueous medium, explored the
possibility of adjusting the pH and solvent composition of the DESI spray to quench the
enzymatic reaction and thus improved the accuracy of the kinetic measurements by preventing
post-ionization reactions.

As an alternative to DESI, matrix-assisted laser desorption ionisation mass spectrometry
imaging (MALDI-MSI) has readily been applied towards the large-scale phenotyping of bacteria
[14, 15]. A related optically-guided MALDI-MS strategy has recently been implemented for the
profiling of microbial colonies for rapid screening of natural product analogue libraries [16].
This impressive development used optical imaging of microbial colonies to direct the laser
coordinates for an automated MALDI-MS screening of approximately 1000 colonies directly
from an imprinted glass slide with an MS sampling rate of about one colony per second.
Reaction products were screened in situ and results overlaid with the optical images;
integration of results allowed for subsequent colony picking and recovery of the desired mutant
strains. The majority of commercially available MALDI-MS instrumentation permit a spatial
resolution of > 100 μm. However, the group of Bernhard Spengler has recently dramatically
pushed this boundary towards much better lateral resolutions down to 1.4 μm [17], thus further
advancing the technique towards single cell resolution and even higher throughput [18].

The coupling of microreactors or continuous flow chemical reactors directly to the mass
spectrometer provides an enhanced ability to characterise unstable reaction products and
reduces the sample volume required (albeit with sufficient mass spectrometer sensitivity). Link
et al. [19] provided a comprehensive example of such an application. They
demonstrated the ability to undertake real-time metabolome profiling by direct
injection of living bacteria, yeast and mammalian cells into a high-resolution mass spectrometer through coupling a peristaltic pump and two six-port valves and automatically sampling from a liquid culture. This approach permitted the automated monitoring of around 300 compounds in 15–30 s cycles over several hours. They investigated the metabolite dynamics in real-time during 2 h starvation and 30 min of growth resumption. The approach suggested that the accumulation of energetically costly metabolites in starved *E. coli* reflects the control strategy to favour cheap metabolic pathways for growth resumption. From an analytical perspective the method permitted real-time metabolome profiling that followed the dynamics of metabolic processes in different organisms over extended periods. The method alleviates retrospective manual sampling, sample preparation and sample manipulation associated with traditional off-line methods.

Progress has also been made on the mass spectrometry techniques available to the synthetic biology community: proton transfer reaction mass spectrometry (PTR-MS) and selected ion flow tube mass spectrometry (SIFT-MS). These techniques are direct injection approaches that utilise chemical ionisation for real-time analysis of volatile organic compounds. PTR-MS has been shown to achieve near-to-real-time monoterpene separation and identification, when coupled to a fast gas chromatography, with sensitivity in the range of 1.2 ppbv from plant material [20]. PTR-MS has also been applied to the real time monitoring of the yeast volatilome [21], detecting more than 300 metabolite features, 70 of which were tentatively identified, in the headspace of *Saccharomyces cerevisiae* cultures over 11 days at 4-h time points. Additional development and application of this technique has been demonstrated by Materic *et al.* [22], who used Selective Reagent Ion PTR-MS to investigate the separation of monoterpene mixtures, which are a particularly common target in recent synthetic biology projects *i.e.* geraniol [23], linalool [24] and limonene [25].

**Global analysis – Metabolomics**

Synthetic biology requires not only the rapid and accurate quantitation of the desired end products; even more important for a systematic engineering of the microbial factories is a thorough understanding of metabolic flux and the regulation of central carbon metabolism to ensure the desired production of target compounds is compatible with maintaining cellular homeostasis and energy balance. Metabolomics, the comprehensive profiling of small molecules in a biological sample, is the obvious method of choice for collecting the necessary data for this kind of analysis, and synthetic biology can build on a continuously refined repertoire of metabolomics approaches [26, 27].

Of the many technological advances in recent years, we only highlight the increasing importance of parallel reaction monitoring (PRM) in metabolomics; the quantitation of intermediates of central carbon metabolism, amino acids and shikimate pathway-related metabolites in engineered strains of *E. coli* [28] is just one important example of its application in synthetic biology. PRM permits the quantitative analysis of multiple targets (237 in this example) [29] with excellent linearity of quantitation, as well as high precision and accuracy. In a related approach, all ion fragmentation acquisition has recently been demonstrated to achieve increased accuracy in metabolite identification for a large number of pre-selected compounds, while at the same time acquiring full scan information to allow the identification of additional metabolites that were initially not targeted [30].

A metabolomics-driven approach was applied to identify non-obvious target genes to further improve the production of 1-butanol [31, 32]. The authors performed quantitative targeted analysis of acyl-CoAs in the CoA-dependent 1-butanol biosynthetic pathway in *Synechococcus elongatus* strains via $^{13}$C-labelling of cell extracts as an internal standard and HPLC-MS analysis. The results indicated several targets for potential improvements of 1-butanol production in
cyanobacteria, such as possible rate-limiting steps (reductive reaction of butanoyl-CoA to butanal) or effective regeneration of free-CoA from butanoyl-CoA to enhance the conversion of pyruvate to acetyl-CoA. In a parallel study addressing 1-butanol production in *E. coli*, the authors examined the metabolomic impact of the deletion of phosphate acetyltransferase, which was performed in an attempt to reduce the amount of acetate produced and simultaneously increase the acetyl-CoA pool. Metabolomics analysis using a targeted ion pair LC-MS/MS method detected a total of 78 metabolites and pointed to several metabolic perturbations caused by the deletion that seemed to be the consequence of a CoA imbalance or insufficient CoA recycling, which caused the undesirable accumulation of side products. Further metabolomics analysis identified the underlying enzymatic bottleneck, alcohol dehydrogenase, and fine-tuning of this activity resolved the CoA imbalance and led to substantially improved 1-butanol titres [31].

A metabolomics approach was also implemented to investigate central metabolism of a fructose repressor (*fruR*) knockout in a recombinant L-tryptophan producing strain of *E. coli* (*E. coli* FB-04) [33]. The authors report more than 80 intracellular metabolites that were altered as a result of the knockout, 23 of which were related to tryptophan biosynthesis. The levels of glycolysis, pentose phosphate and TCA cycle intermediates were consistently increased, and levels of shikimate derivatives (direct tryptophan precursors) and L-glutamine were decreased in the knockout strain, which also showed a substantially increased tryptophan production. The interpretation of these results illustrates very clearly the pitfalls of using steady-state metabolome profile information as a proxy for metabolic fluxes, which are of central interest for synthetic biology: based on increased levels of glycolytic and pentose phosphate pathway intermediates, the authors conclude that the *fruR* knockout enhanced metabolic flow through these two pathways which provide the substrates for L-tryptophan biosynthesis. However, the TCA cycle, which directly competes with tryptophan biosynthesis shows an equally increased level of its intermediates, and the only pathway for which direct flux measurements are available, tryptophan biosynthesis itself, shows a consistent decrease in its key intermediates, despite an increase in flux by 62.5% (from 0.024 to 0.039 g/L/h).

A subsequent study combining metabolomics and 13C fluxomics provided more detailed insights into the metabolic flux redistribution in an *E. coli* strain overproducing shikimic acid with high titres and yields: Rodriguez et al. [34] used an engineered AR36 *E. coli* strain constitutively expressing six proteins encoded in a synthetic operon promoting high-yield production of shikimic acid from glucose. Comparative metabolomics of a production strain and parental strains (carrying either no plasmid or “empty plasmid”) was used to track the levels of seven exometabolites and 25 endometabolites over time. It revealed a global remodelling of carbon and energy metabolism in the high producer. This resulted in reduced carbon available for oxidative and fermentative pathways and increased levels of endometabolites involved in energy pathways, preventing the depletion of essential intermediates, such as PEP and ATP. Both glycolytic flux and TCA cycle activity were substantially reduced in this overproduction scenario (43 g/L of shikimate in 30 h on complex medium).

Given its importance as a provider of essential precursors for a diverse range of biotechnologically important biochemicals, it is not surprising that the shikimate pathway has been the target of dedicated metabolomics method development: e.g., Lai et al. [35] contributed a robust HPLC method for the quantification of aromatic substrates, products and pathway intermediates in order to accelerate strain engineering for industrial production of aromatics as biosynthetic molecules. The achieved limits of detection between $10^{-10}$ – $10^{-13}$ mol make the method suitable for endometabolome and exometabolome analysis of engineered strains.

Another example of a metabolomics-based strategy for strain engineering (this time utilising a GC-MS analytic platform) is the study by Teoh et al. [36] investigating phenotypic differences in
growth rates and metabolite profiles of nineteen single-deletion *S. cerevisiae* mutant strains cultivated under stress-free and under 1-butanol stress conditions (growth inhibition caused by higher alcohols (e.g. 1-butanol) is considered as a bottleneck in their biosynthetic production). Metabolites associated with improved growth rates under stress conditions were identified, and new stress-resistant mutant yeast strains were successfully predicted based on their metabolite profiles. This approach illustrates the potential of metabolomics as a predictive screening tool to inform semi-rational strain engineering approaches.

Finally, metabolomics has been applied for the monitoring of isoprenoid precursors production, another classic target for synthetic biology [37, 38]. In a study by Kirby et al. [39], who report for the first time the functional expression of an extensively engineered functional 1-deoxy-D-xylulose 5-phosphate (DXP) pathway in *S. cerevisiae* which normally utilizes the mevalonate pathway, which has a lower theoretical yield. Metabolite-guided DXP pathway balancing, by LC-MS quantification of intermediates in cultures exhibiting various levels of flux, appeared to be a successful approach for identifying a bottleneck in the pathway. An engineered strain exclusively using the DXP pathway achieved an endpoint biomass 80% of that of the same strain using the mevalonate pathway under low aeration conditions.

**Molecular networking – moving forward**

The main challenge of untargeted metabolomics is compound annotation; the persistent difficulties of confidently identifying the detected metabolites currently seriously limits the utility of the MS data acquired. Molecular networking, a visualisation method for tandem MS data, is a powerful complement to traditional de-replication methods [40]. This approach allows for the detection of sets of spectra from related molecules (“spectral networks”), even in the cases when these spectra are not matched to any known compounds. The approach is based on the assumption that similar molecules have similar MS fragmentation patterns so they will tend to cluster closely within a network. Each spectrum (ideally derived from a single compound) is visualised as a network node, and the edges between nodes represent a degree of similarity between spectra. The thicker the line, the more MS/MS fragment ions are shared by the two connected nodes. Nodes can be supplemented by such information as a compounds abundance, biochemical activity, origin etc. Molecular networking led to the development of Global Natural Products Social Molecular Networking (GNPS), a metabolomic data-driven platform for the storage, sharing, analysis, and knowledge dissemination of tandem MS spectra where one is able to annotate natural product data via continuous de-replication. [41].

Although improvements are still required to obtain unambiguous analysis of molecular networks such as efficient integration with existing LC-MS detection strategies, enhancement of pre-processing and universal optimal acquisition methods [40, 42, 43], its applications are expanding fast [44-49] and it will soon become an indispensable metabolomics tool in exploratory analyses for the synthetic biology of novel natural products. For example, in a recent study Crüseman and colleagues [46] screened 146 marine *Salinispora* and *Streptomyces* strains using HPLC-MS/MS, molecular networking, and the Global Natural Products Social (GNPS)[41] platform and explored the impact of differing culturing and extraction techniques. The systematic investigation of the effect of these parameters clearly demonstrated how much inherent chemical diversity could be missed when just one culture and extraction protocol is utilised to assess metabolic capacity. This example demonstrated how the application of molecular networking permits the rapid optimisation of experimental parameters that can subsequently be implemented early in the discovery workflow.

Okada et al. [50] used molecular networking for the investigation of the influence of trimethoprim (Tmp) antibiotic on the secreted metabolome of *Burkholderia thailandensis* E264. The untargeted comparison of Tmp-induced and uninduced samples (utilising HPLC-QToF-
MS/MS) resulted in ~240 metabolites of interest (with >100 compounds observed only for the induced samples). Organising them into 14 sub-networks followed by NMR analysis enabled rapid identification of 40 compounds including analogues of known compounds and a group of new molecules, acybolins, showing that molecular networking aids rapid identification of compounds compared to traditional workflows.

In related work, von Eckardstein et al. [47] used bioactivity-guided untargeted LC-MS/MS analysis and molecular networking in the search of new antibiotic agents from Xanthomonas albilineans. Over 20,000 MS/MS spectra acquired from crude extracts and bioactive fractions were organised into a molecular network via the GNPS portal, which allowed for the identification of potential derivatives in the albicidin sub-network. The group reported eight new natural albicidin derivatives with unambiguous identification.

**Conclusions**

There are still some challenges to overcome, but both synthetic biology and metabolomics are very dynamic fields that are forging an ever-closer alliance. An example that illustrates the integral role of metabolomics in synthetic biology pipelines is the recently published multi-omics workflow to characterise strain variation in engineered *E. coli* [51]. It is certain that in coming years we will see a rapid deepening of the technical and conceptual integration of metabolic profiling methods within the Design – Build – Test cycle, and in particular the emergence of additional tools to facilitate the flow of data and insights between the analytical machinery (Test) and its users in the Design and Build stages of strain engineering.

**Acknowledgements**

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**References and recommended reading**

Papers of particular interest, published within the period of review, have been highlighted as:

• of special interest

•• of outstanding interest


In this article, the authors show how a optically-guided MALDI-MS allows the rapid identification and recovery of desirable strains from large populations of microbial colonies.


This study illustrates the high-throughput on-line monitoring of the volatile metabolome of microbial colonies enabled by automatic proton transfer reaction mass spectrometry.


Here, the authors present the application of parallel reaction monitoring in metabolomics, as a method to rapidly characterize the metabolic pathway reorganization observed in engineered *E. coli* strains.


Using global metabolic profiling, the authors identify a non-obvious new target for enhancing the production of a desired end compound in engineered microbes.


A combination of label-free global metabolite profiling and stable-isotope labelled fluxomics revealed new potential rate-limiting steps in a well-studied metabolic pathway.


A combination of metabolomics profiling and fluxomics experiments provided detailed insights into the profound metabolic consequences of the overproduction of a key precursor for an important family of natural products.


This study illustrates the power of computational metabolomics, introducing a new algorithm for the dereplication of natural products libraries using molecular networking strategies.


Metabolite networking enabled the rapid parallel exploration of the secondary metabolome of more than two hundred ecologically diverse strains of microbes, facilitating dereplication, while at the same time identifying structural relationships between newly discovered compounds.


<table>
<thead>
<tr>
<th>Technique/Approach</th>
<th>Full Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Metabolomics</td>
<td>—</td>
<td>The untargeted, non-biased detection and identification of all low-molecular weight compounds (metabolites) present within a biological sample or system.</td>
</tr>
<tr>
<td>MS</td>
<td>Mass Spectrometry</td>
<td>Analytical technique based on the ionisation of analytes (e.g., by DESI, MALDI, PTR or SIFT; see below), the subsequent separation of ions according to mass/charge ratio, and their detection and quantification.</td>
</tr>
<tr>
<td>MSI</td>
<td>Mass Spectrometry Imaging</td>
<td>Mass spectrometry is conducted in a spatial manner thus permitting the visualisation of the two-dimensional localisation of analytes within a sample, for example across a microbial colony growing on an agar plate.</td>
</tr>
<tr>
<td>DESI-MS</td>
<td>Desorption Electrospray Ionization Mass Spectrometry</td>
<td>Ambient ionization technique using a nebulized electrospray. Highly charged microdroplets collect analytes from the surface of the sample prior to secondary droplets carrying the analyte to the MS. This ionization technique is particularly suitable for MSI.</td>
</tr>
<tr>
<td>IM-MS</td>
<td>Ion Mobility Mass Spectrometry</td>
<td>A variant of MS, with additional separation of ions according to the time it takes for them to travel through a drift tube with a homogeneous, continuous electric field in the presence of a neutral gas. This leads to separation of ions according to size and shape (collision cross section), complementing the mass/charge information available in traditional MS.</td>
</tr>
<tr>
<td>MALDI-MS</td>
<td>Matrix Assisted Laser Desorption Ionization Mass Spectrometry</td>
<td>Ionization approach whereby a matrix (an energy-absorbing small organic compound) is applied to/mixed with a sample. A laser applied to the matrix:sample mix excites the matrix molecules and leads to the generation of volatilized ions which subsequently enter the MS. This technique is suitable for MSI.</td>
</tr>
<tr>
<td>PTR-MS</td>
<td>Proton Transfer Reaction Mass</td>
<td>A soft ionization technique using an ion beam of protonated water</td>
</tr>
<tr>
<td>Method</td>
<td>Technique</td>
<td>Description</td>
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<td>----------------------</td>
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<tr>
<td>Spectrometry</td>
<td></td>
<td>molecules, H$_3$O$^+$, as an ion source to protonate (and thus ionize) volatile analytes. This technique permits for real-time monitoring of organic molecules in the gas phase.</td>
</tr>
<tr>
<td>SIFT-MS</td>
<td>Selected-Ion Flow-Tube Mass Spectrometry</td>
<td>Similar to PTR-MS, this soft ionisation technique uses precursor ions in the gas phase to ionize volatile analytes. The precursor ions are generated by a microwave plasma ion source, and a single ion species can be selected (H$_3$O$^+$, NO$^+$ or O$_2^-$) to perform as reactant ion. Neutral volatile analyte molecules react with the precursor ions and undergo ionization. This technique permits for real-time monitoring in the gas phase.</td>
</tr>
<tr>
<td>Molecular Networking</td>
<td>—</td>
<td>A computational method for MS data analysis that allows for the identification of sets of spectra from chemically related molecules (“spectral networks”), based on similarities in molecular fragmentation patterns, even in the cases when the spectra are not matched to any known compounds.</td>
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</tbody>
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