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Simultaneous quantification of multiple bacterial metabolites using surface-enhanced Raman scattering

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Given the commercial importance of the compounds produced by genetically modified organisms, there is a need for screening methods which facilitate the evaluation of newly developed strains, especially during the phase of proof-of-concept development. We report a time-efficient analysis method for the screening of bacterial strains, which enables the detection of two structurally similar secondary bacterial metabolites. By combining liquid-liquid extraction and surface-enhanced Raman scattering we were able to quantify p-coumaric acid and cinnamic acid, produced by genetically modified E. coli from tyrosine and phenylalanine, respectively. With the simple sample pre-treatment method, and by applying a partial least squares data analysis method, we simultaneously detected the analytes from four E. coli strains cultured in the presence or absence of tyrosine and phenylalanine.

Introduction

Metabolic engineering enables the direct modulation of cellular activities by manipulation of enzymatic, transport and regulatory functions, for metabolite overproduction or improvement of cellular properties.1 This field gained importance for production of food additives, biofuels and pharmaceuticals, focusing both on the optimization of existing processes and on the development of new ones.2 The use of genetically modified organisms is particularly convenient for the production of compounds which are too complex to be chemically synthetized, or too time- and resource-consuming to be extracted from natural sources.3 Additionally, although in most cases the production is focused on a single molecule, one organism can be modified to synthetize more than one analyte simultaneously.4,5 When developing a proof-of-concept microbial strain, several genetic modifications are needed to boost the desired process and prevent the competing ones, due to the complexity of microbial metabolism.6 Therefore, a large number of strains must be screened to identify the best performing ones, with repeated optimization cycles resulting in a costly and time-consuming process, lasting up to several years.2 Bioinformatic tools and new technologies have contributed to fast design and modeling of metabolic pathways, and to rapid genomic modification. Nowadays genetic engineering can be performed in massive parallel reactions, resulting in either specific clones or in libraries of different cells.7 DNA synthesis for direct use or in gene assembly8,9 and techniques such as mutagenesis by UV radiation are also capable of generating myriads of variation.10 When constructing genetic manipulations, specific gene insertions can be done in in less than 20 minutes,11 however, getting data and selection of the best producing strain is still time consuming.7,12 Therefore, there is still a need for fast and high-throughput screening techniques to speed up testing, which is a crucial step and it has been described as the bottleneck of the entire development process.7,12

The analytical techniques that are commonly used for analysis (e.g. gas chromatography, high-performance liquid chromatography), are accurate and robust, but they require expensive instrumentation operated by skilled personnel, time-consuming optimization of protocols and long separation procedures.12 Raman spectroscopy, a non-destructive, fast and label-free technique, is suitable for collecting molecule-specific information from samples, even in small quantities.13 In spite of the advantages offered by Raman spectroscopy, detection of molecules with a small Raman cross-section or at low concentrations in complex media is often challenging. To be able to detect in the aforementioned conditions, high laser power and long acquisition time are required, which could result in the degradation of the analyzed molecules. Additionally, fluorescence can interfere with the Raman signal.14 On the other hand, surface-enhanced Raman scattering (SERS) enables detection at low concentration,15 or even of single molecules,16 due to its ability to enhance the Raman signal of a molecule with reported enhancement factors up to 1014, through the use of metallic nanostructured...
Materials and methods

Chemicals, E. coli cultures and HPLC detection

Stock solutions of 100 mM pHCA and 100 mM CA were freshly prepared in EtOH 99%. For SERS calibration standards, pHCA and CA were diluted in dichloromethane (DCM) or in control supernatant (obtained from a non-pHCA/CA producing E. coli strain (CBI786)). In addition, for SERS and HPLC experiments the analytes were diluted in M9 medium. HCl 32% was used for acidification of samples and DCM as the organic phase for LLE. Stock solutions of 50 mM Phe and 50 mM Tyr were prepared in water and NaOH at pH ~ 14 respectively. E. coli strains (CBI800, CBI786, CBJ951, CBJ789), expressing TAL and PAL-encoding genes from IPTG-inducible promoters4 (Fig. 1), were grown in M9 medium with 1% glucose, 2 mM Tyr and/or 2 mM Phe, 1 mM IPTG and antibiotics for maintenance of plasmids for 22 h as described in our previous work.32 For quantification of pHCA and CA produced by E. coli, bacterial supernatant samples were obtained from each strain by centrifugation (10 min at 10000 g, 4 °C), and filtration through 0.2 µm filters. The concentration of pHCA and CA in samples was measured with reversed-phase HPLC by separation on a HS-F5 column (Sigma-Aldrich, St. Louis, MO, USA) with previously described mobile phases (ammonium formate buffer and acetoni-trile), with an overall analysis time of approximately 15 min per sample.32 The absorbance was measured at 333 nm for pHCA and at 277 nm for CA.4 Aqueous solutions were prepared with ultrapure water obtained from a Milli-Q purification system (Millipore Corporation, Billerica, MA, U.S.), and all the chemicals were purchased from Sigma-Aldrich (St. Louis, MO, USA).

SERS chip fabrication

The gold-capped silicon nanopillar substrates were fabricated as described by Wu et al.,31 with 4 min etching time, followed by 1 min O2 plasma cleaning and deposition of 220 nm Au at a rate of 10 Å/s. Wafers were diced with a laser micromachining tool (3D-Micromac AG, D-09126 Chemnitz, Germany) to create 4x4 mm2 chips used for measurements.

SERS data acquisition and analysis

SERS sensing was performed on 4x4 mm2 substrates using 5 µL DCM or 2 µL EtOH droplets after the samples were completely dried. The measurements were performed with a DXRxi Raman Imaging Microscope (Thermo Fisher Scientific Inc., Waltham, MA, US). The optical microscope was coupled to a spectrometer 5 cm−1 FWHM and ±2 cm−1 wavenumber accuracy. SERS spectra were collected as previously described at 780 nm wavelength with a laser power of 1 mW, 10x objective lens, 50 µm slit and 3.6 µm diameter estimated laser spot. Maps of 48 points with a 100 µm collection step were collected on the surface of each chip, and the spectrum collected in each point was averaged over 3 acquisitions of 0.05 s each, resulting in a total detection time of approximately 3 min per sample.
Data pre-processing and quantitative analysis were performed with MatLab (version 8.4, MathWorks, Natick, MA, US) and TQ Analyst (version 9.2, Thermo Fisher Scientific Inc., Waltham, MA, US). An average SERS spectrum was obtained for each map after polynomial baseline correction (7th order) and outlier removal, followed by the application of a partial least squares (PLS) algorithm to build a quantitative model. Standards with known concentrations were used to build the calibration models, and validation samples were randomly chosen for each concentration within the dataset in each case. The models were developed based on a first derivative algorithm in the wavenumber range 800 - 1700 cm⁻¹, in conjunction with quadratic baseline removal.

Sample extraction

The experimental procedure for LLE, based on Krygier et al.⁵² is presented in Fig. 2. The extraction was carried out as previously described by Morelli et al.³² using 500 µL of bacterial supernatant or spiked M9 medium containing pHCA, CA and/or Tyr and Phe. The sample was acidified with 50 µL 32% HCl(aq) (Fig. 2a) and vortexed for optimal mixing. 500 µL DCM was added to the sample (Fig. 2b) and vortexed again, followed by 30 min static incubation at room temperature. The aqueous phase was then removed and diluted 10 times with phosphate buffer (pH 8, 0.04 M) for HPLC analysis (Fig. 2c), in order to increase the pH to 6, whereas the organic phase was directly used for SERS analysis (Fig. 2d and e).

Results and discussion

LLE and SERS for extraction and detection of pHCA and CA

As described in our previous works,³¹,³²,³⁴ genetically modified E. coli strains are cultured in a growth medium containing appropriate salts, nutrients and antibiotics, with substrates (e.g. Tyr, Phe) at starting concentrations up to 2 mM. According to the genes expressed by each strain, and according to their synthesis efficiency, substrates can be found at high concentrations in bacterial supernatant. The SERS spectra of Tyr and Phe in EtOH, highlighted in red and black respectively in Fig. 3a, partially overlap with the signal of pHCA and CA (purple and blue graphs). For instance, both CA and Phe present a peak at 1000 cm⁻¹, whereas overlapping peaks can be found in the region 1100 – 1300 cm⁻¹ for Tyr, pHCA and CA. Therefore, all the molecules contribute to the SERS spectrum of the mixture (Fig. 3a, green) to a different extent.

LLE, a common extraction method for phenolic compounds,⁵³–⁵⁵ was used for separation of pHCA and CA, according to the experimental procedure depicted in Fig. 2. LLE enables the partition of a compound between two immiscible phases. In general, uncharged compounds tend to be more soluble in organic solvents, whereas charged molecules tend to stay in the aqueous phase. By regulating the pH of the aqueous sample, uncharged molecules, such as pHCA and CA, can be extracted to the organic phase whereas charged molecules (e.g. Tyr, Phe and salts present in the growth medium) remain in the aqueous phase.

In our previous works, we combined LLE and the here exploited gold-capped silicon nanopillar substrates for extraction and SERS detection of pHCA.³²,³³ Besides excluding Tyr and salts, measuring the target analyte in the organic phase (DCM) provided the optimal working conditions for metal-capped nanopillar substrates, since it was demonstrated that wetting the substrate with an organic solvent improves the leaning of silicon nanopillars,³⁶ leading to a better SERS signal enhancement.

With LLE we were able to extract both pHCA and CA from aqueous samples. The effective exclusion of Tyr and Phe was demonstrated by performing LLE on medium spiked with Tyr and Phe only (Fig. 3b, light blue). It can be observed that the obtained signal was comparable to the one recorded from the control (Fig. 3b, orange). Additionally, when LLE was not performed, no visible vibrational peaks could be recorded, even after diluting the sample with EtOH (Fig. 3b, brown).³²,³⁴

In Fig. 3b, the green spectrum represents the signal obtained after LLE of a sample containing pHCA, CA, Tyr and Phe. Vibrational peaks originating from both pHCA (e.g. at 1169 cm⁻¹) and CA (e.g. at 1000 cm⁻¹) could be observed, as also shown in Fig. 3a, without the influence of Tyr and Phe. Therefore, LLE enabled the exclusion of both salts and substrates with interfering vibrational peaks.

Quantification of pHCA and CA spiked in M9 medium

As previously described, LLE enables the partition of a compound between two immiscible liquid phases, according to the affinity of the compound towards each phase. The partition of pHCA and CA in this work was evaluated through the extraction efficiency (EE), calculated as:

\[
\% \, EE = \left( \frac{[S_i^2]}{[S_i]} \right) \times 100
\]

where \([S_i]\) is the initial concentration, known in spiked samples, and \([S_i]^2\) is the concentration of \(S\) in the organic phase after extraction.

Based on our previous study,³² we found that EE was 9.8 ± 1.1 % for pHCA. However, in the case of simultaneous detection of pHCA and CA, it was important to evaluate the EE of CA as well as of both analytes in solution.

Samples of M9 medium were used, spiked with 500 µM of pHCA and CA in different proportions (pHCA/CA: 500/0, 400/100, 300/200, 200/300, 100/400, 0/500 µM); the concentration range was similar to the one expected in bacterial supernatant (see Fig. 5c). After LLE, the aqueous phase was removed, diluted 10 times with phosphate buffer (pH 8) and measured with HPLC in order to calculate the concentration of pHCA and CA left in aqueous phase, as depicted in Fig. 2. The EE of pHCA was confirmed to be 9.4 ±
pHCA concentration (RMSEC = 5.65 µM, r2 = 0.993, LOD = 90 µM), and CA (RMSEC = 28.1 µM, r2 = 0.992 and RMSEP = 29.8 µM, r2 = 0.992, LOD = 70 µM).

As shown in Fig. S2, the model was able to separate the signal of the analytes from the background. However, the influence of the background signal recorded from supernatant controls was more significant than in the case of M9 controls, and the analyte spectra had a few differences compared to pHCA and CA SERS spectra in EtOH. For instance, we found a prominent peak originating from the control samples at 1000 cm⁻¹, therefore the overlapping characteristic CA peak could not be used for CA quantification. Nevertheless, significant spectral features were successfully identified for both pHCA and CA, such as the pHCA peak at 1169 cm⁻¹, and the CA peak proportion in the 1580 – 1660 cm⁻¹ spectral region.

Bacterial supernatant samples were obtained from E. coli strains CBJ786, CBJ951, CBJ800 and CBJ789, cultured as described in Materials and Methods. According to the presence of Tyr and/or Phe in the growth medium (indicated in Fig. 5c by T and/or P next to the number of the corresponding strain), different concentrations of pHCA and CA were obtained. Supernatant samples were processed with LLE and the extracts were measured with SERS (Fig. S3), applying the calibration models in Fig. 5a and b for quantification. With this method, as shown in Fig. 5c, pHCA and CA were successfully quantified in real supernatant samples, with results in good agreement with HPLC.

The pre-treatment time was 35 min for all the samples, plus 3 min for SERS acquisition of each sample, versus 15 min for each HPLC run, with a significant gain in time-to-result. Moreover, the detected pHCA and CA concentrations were within or below ranges reported in several studies, proving that the developed method is suitable for differentiating engineered microbial strains overproducing pHCA and CA.

Conclusions

In this work we demonstrated the simultaneous extraction and quantification of bacterial metabolites in E. coli supernatant. The advantages of the previously developed method, based on LLE sample pre-treatment and SERS acquisition, were confirmed and used in combination with a simple PLS model for quantification of multiple analyte mixtures in real supernatant samples. The result was a simple and effective method for fast and quantitative screening of bacterial strains, based on the amount of produced metabolites. The presented work demonstrates the applicability of the method during the development of a proof-of-concept strain, where there is a need for a time- and cost-efficient screening of a large number of strains.

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**Conflict of interest**

There are no conflicts of interest to declare.

**References**


Figures

Fig. 1 TAL and PAL enzymatic reactions.
Fig. 2 Experimental procedure for extraction of pHCA/CA from *E. coli* supernatant or spiked M9 medium samples. (a) The samples were acidified at pH 0.1 and (b) DCM was added to enable liquid-liquid extraction (inset). (c) The concentration of pHCA and CA in aqueous phase after extraction was analyzed with HPLC, whereas (d) droplets of organic phase were poured on SERS substrates, dried and (e) collected with a Raman microscope.
Fig. 3 (a) SERS spectra of 500 µM pHCA, CA, Tyr and Phe in EtOH 99% (green) and of 500 µM of each analyte, after background subtraction. Black and red dots highlight the vibrational peaks of Phe and Tyr respectively, which can also be found in CA and pHCA spectra. (b) SERS spectra of M9 medium spiked with 500 µM pHCA, CA, Tyr and Phe 2-fold diluted with EtOH 99% (brown) prior to liquid-liquid extraction (LLE), and of organic extracts of medium spiked with the 4 analytes (green), with only Tyr and Phe (light blue), and with no analytes (orange). The spectral features of pHCA and CA, overlapped in the mixture, are highlighted with purple and blue dots respectively. The spectra were shifted with arbitrary offsets for ease of representation.
Fig. 4 PLS calibration model in DCM for (a) pHCA and (b) CA, with each point representing a SERS map (data collected on triplicate chips with 2 maps on each chip, n = 6). (c and d) Quantification of the mixture spiked in M9. SERS data were collected on triplicate chips, with two maps on each chip, with error bars representing standard deviation (n = 6).
**Fig. 5** PLS calibration model of (a) pHCA and (b) CA in spiked supernatant samples pre-treated with LLE, with each point representing a SERS map (data collected on triplicate chips with 2 maps on each chip, n = 6). (c) SERS quantification of real samples compared with HPLC data. T and P indicate Tyr and/or Phe enriched growth medium for *E. coli* pHCA and CA production. SERS data were collected on triplicate chips, with two maps on each chip (n = 6), whereas HPLC data were averaged over triplicate injections (n = 3).