



Intra-Ramanome Correlation Analysis Unveils Metabolite Conversion Network from an Isogenic Population of Cells

Yuehui He,^{a,b,c} Shi Huang,^{a,b,c,d} Peng Zhang,^{a,b,c} Yuetong Ji,^{a,b,c}  Jian Xu^{a,b,c}

^aSingle-Cell Center, CAS Key Laboratory of Biofuels, Shandong Key Laboratory of Energy Genetics and Shandong Energy Institute, Qingdao Institute of BioEnergy and Bioprocess Technology, Chinese Academy of Sciences, Qingdao, Shandong, China

^bLaboratory for Marine Biology and Biotechnology, Qingdao National Laboratory for Marine Science and Technology, Qingdao, Shandong, China

^cUniversity of Chinese Academy of Sciences, Beijing, China

^dDepartment of Pediatrics and Center for Microbiome Innovation at Jacobs School of Engineering, University of California San Diego, La Jolla, California, USA

ABSTRACT To reveal the dynamic features of cellular systems, such as the correlation among phenotypes, a time or condition series set of samples is typically required. Here, we propose intra-ramanome correlation analysis (IRCA) to achieve this goal from just one snapshot of an isogenic population, via pairwise correlation among the cells of the thousands of Raman peaks in single-cell Raman spectra (SCRS), i.e., by taking advantage of the intrinsic metabolic heterogeneity among individual cells. For example, IRCA of *Chlamydomonas reinhardtii* under nitrogen depletion revealed metabolite conversions at each time point plus their temporal dynamics, such as protein-to-starch conversion followed by starch-to-triacylglycerol (TAG) conversion, and conversion of membrane lipids to TAG. Such among-cell correlations in SCRS vanished when the starch-biosynthesis pathway was knocked out yet were fully restored by genetic complementation. Extension of IRCA to 64 microalgal, fungal, and bacterial ramanomes suggests the IRCA-derived metabolite conversion network as an intrinsic metabolic signature of isogenic cellular population that is reliable, species-resolved, and state-sensitive. The high-throughput, low cost, excellent scalability, and general extendibility of IRCA suggest its broad applications.

IMPORTANCE Each isogenic population of cells is characterized by many phenotypes, which change with time and condition. Correlations among such phenotypes are fundamental to system function, yet revelation of such links typically requires multiple samples. Here, we showed that, by exploiting the intrinsic metabolic heterogeneity among individual cells, such interphenotype correlations can be unveiled via just one snapshot of an isogenic cellular population. Specifically, a network of potential metabolite conversions can be reconstructed using intra-ramanome correlation analysis (IRCA), by pairwise correlation of the thousands of Raman peaks or combination of peaks among single-cell Raman spectra sampled from just one instance of the cellular population. The ability to rapidly and noninvasively reveal intermetabolite conversions from just one snapshot of one sample should usher in many new opportunities in functional profiling of cellular systems.

KEYWORDS ramanome, intra-ramanome correlation analysis (IRCA), intra-ramanome correlation network (IRCN), single-cell Raman spectroscopy, phenotypic heterogeneity

Each isogenic population of cells is characterized by many phenotypes that change with time and condition. Such phenotypes can be related to cellular metabolism, such as the amount of a particular metabolite produced under a given time and condition. Links among such metabolism-related phenotypes are a fundamental property that underlies proper functioning of cellular systems (1–3). For example, correlation in abundance among metabolites can unravel, via metabolome-wide association studies

Citation He Y, Huang S, Zhang P, Ji Y, Xu J. 2021. Intra-ramanome correlation analysis unveils metabolite conversion network from an isogenic population of cells. *mBio* 12:e01470-21. <https://doi.org/10.1128/mBio.01470-21>.

Editor Sang Yup Lee, Korea Advanced Institute of Science and Technology

Copyright © 2021 He et al. This is an open-access article distributed under the terms of the [Creative Commons Attribution 4.0 International license](https://creativecommons.org/licenses/by/4.0/).

Address correspondence to Jian Xu, xujian@qibebt.ac.cn.

Received 16 June 2021

Accepted 5 August 2021

Published 31 August 2021

(MWAS) of metabolomic data sets from high-resolution mass spectrometry (MS), the links among target metabolites that characterize a certain disease (4–6). Therefore, strategies and methods for rapidly and comprehensively detecting and profiling such links are of interest.

To detect such links, variations in metabolism-related phenotypes are required. As a result, typically, a time series or condition series set of samples is characterized for metabolite contents first and then correlated across the samples, with the resolution of detection generally dependent on sample size (7–9). For example, in order to understand the mechanism of oil production in microalgae, the lipidomes of an industrial microalga, *Nannochloropsis oceanica*, were measured in triplicate by electrospray ionization MS over eight time points (3, 4, 6, 12, 24, 48, 72, and 96 h) under nitrogen-replete (N+) or nitrogen-depleted (N-) conditions; then the profiles were correlated across the 48 samples, which revealed several temporal patterns among the triacylglycerol (TAG) species that suggest distinct regulation mechanisms; e.g., TAGs with lower degrees of desaturation were induced at the early stages of N-, while those containing polyunsaturated fatty acids (PUFAs) increased considerably only at later stages (10).

However, within a single sample of an isogenic population of cells, at any given time or condition, variations in metabolism-related phenotypes among individual cells are inherent and universal, due to the stochasticity of gene expression in cells (11, 12). Therefore, an intriguing question is, can such phenotypic variations among individual cells, instead of those among multiple samples, be exploited to predict the links among metabolism-related phenotypes? Specifically, there are three hypotheses: (i) every cell can be quantitatively profiled for its various metabolism-related phenotypes as an independent biological replicate; (ii) intercellular variations of the phenotypes can be measured rapidly and simultaneously for many cells; (iii) intercellular correlation of the phenotypes can unveil important functional features of the system.

To test these hypotheses, we employed single-cell Raman microspectrometry, which captures the *in vivo* chemical profiles of a cell in a rapid, label-free, and nondestructive manner (13–15). In a single-cell Raman spectrum (SCRS) (13–15), each of (or a combination of) its thousands of Raman peaks potentially represents a specific metabolism-related phenotype, such as the presence and the concentration of a metabolite synthesized by the cell (16). Therefore, just like a portrait can reveal multiple facial features from a human individual, an SCRS can unveil cellular phenotypes (i.e., functions) in a “landscape-like” manner, i.e., simultaneously revealing multiple metabolism-related phenotypes of the cell at that particular state (16). For example, in a single microalgal cell (e.g., *Chlamydomonas reinhardtii* and *N. oceanica*), starch content can be quantified by the specific Raman peaks of 478 cm^{-1} (C-C-C deformation) and 940 cm^{-1} (C-O stretching; C-O-C and C-O-H deformation; α -helix C-C backbone), while the content of triacylglycerol (TAG) can be modeled by $1,441\text{ cm}^{-1}$ (alkyl C-H₂ bend) and $2,851\text{ cm}^{-1}$ (C-H₂; C-H₃ asymmetric and symmetric stretches) (17, 18). In addition, the degree of lipid unsaturation in a cell can be measured by $I_{1,658}/I_{1,441}$, i.e., the ratio of $1,658\text{ cm}^{-1}$ (allyl C=C stretches which are proportional to the amount of unsaturated C=C bonds) and $1,441\text{ cm}^{-1}$ (alkyl C-H₂ bends which are proportional to the amount of saturated C-C bonds) (18, 19). On the other hand, based on the full spectra of SCRS, the contents of many compounds in a cell can be derived, such as protein, starch, and TAG in *C. reinhardtii* (18) and astaxanthin, and β -carotene and chlorophyll in *Haematococcus pluvialis* (20), via chemometric multivariate methods including partial least square regression (PLSR) and multivariate curve resolution (MCR).

We previously proposed the “ramanome” concept (16, 21). A ramanome is the collection of SCRS (one from each cell) randomly sampled from a given instance of an isogenic cellular population and thus represents a single-cell-resolution metabolic snapshot of the population (16, 21). Here, by treating every SCRS in a ramanome as one biological replicate and each of its peaks or combination of peaks as a metabolism-related phenotype, we established a formal framework called intra-ramanome correlation analysis (IRCA; Fig. 1). From just one single ramanome, by pairwise correlating all the Raman peaks of SCRS among the individual cells, IRCA unveils a network of

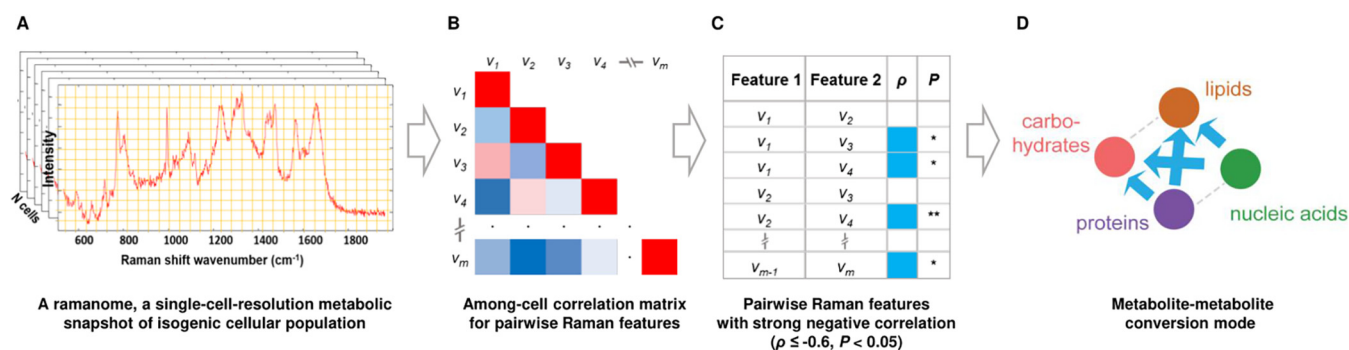


FIG 1 The principle and workflow of intra-Ramanome correlation analysis (IRCA).

potential metabolite conversions. Based on a series of Ramanome-profiling experiments on *C. reinhardtii* mutants and additional microalgae, fungi, and bacteria, we showed that IRCA is a rapid, low-cost, high-throughput, landscape-like, and universally applicable method for unraveling metabolic features of cellular systems.

RESULTS

IRCA predicts the interconversion among starch, protein, and TAG from one single instance of an isogenic microalgal population. To test the IRCA concept (Fig. 1), we employed the unicellular microalga of *C. reinhardtii* CC124 as the first model. Triplicate isogenic cultures of CC124 grown under nitrogen depletion (N-) were sampled from 16 time points from 0 h to 8 days (i.e., 48 Ramanomes in total; see Fig. S1; Materials and Methods). For cellular protein, starch, and TAG contents, each sample was analyzed via two strategies: (i) at the population level, via various conventional techniques which all require metabolite extraction from cell lysates (bicinchoninic acid [BCA] protein assay kit, amyloglucosidase/ α -amylase method and thin-layer chromatography plus gas chromatography/mass spectrometry [TLC-GC-MS]; Fig. 2A to C; also see Text S1) and (ii) at the single-cell level, via full-SCRS-based quantitative partial least square regression (PLSR) models for metabolite contents, which are noninvasive and label-free (18) (~20 randomly selected cells per sample and thus 60 per time point; Fig. 2D to F). The PLSR model for starch content, for example, was established using the population-level measurements (i.e., via the amyloglucosidase/ α -amylase method; see Text S1) and the averaged SCRS of 20 cells in the corresponding biological replicate (i.e., one Ramanome). The correlation coefficients (R^2) of protein, starch, and TAG contents between the two strategies above are 0.9924, 0.9892, and 0.9686, respectively, confirming high accuracy of simultaneous quantification of protein, starch, and TAG for a *C. reinhardtii* cell via its full SCRS (18). Notably, for each phenotype, at any instance of the population, the degree of intercellular heterogeneity is high, and can vary greatly along the 8 days (see Fig. S1G to I). Specifically, for protein, both minimal and maximal single-cell contents in the population were decreasing with time (see Fig. S1G), while for starch, the trends are exactly the opposite (see Fig. S1H). As for TAG, the maximal content was increasing, while the minimal content remained at the baseline (i.e., a subpopulation of non-TAG-producing cells was always present), with the degree of within-population heterogeneity continuing to grow (i.e., the “delta” in Fig. S1I).

Notably, at the population level, significant correlation (defined by the Pearson correlation coefficient, or ρ) was observed in each of three phenotype pairs among the 48 populations—protein-starch ($\rho = -0.926$; Fig. 2A), starch-TAG ($\rho = 0.639$; Fig. 2B), and protein-TAG ($\rho = -0.848$; Fig. 2C). This indicates protein-starch conversion, protein-TAG conversion, and accordant change of starch and TAG contents (22, 23). Intriguingly, at the single-cell level, correlation for the 960 cells collectively reached a consistent conclusion, albeit with lower correlation coefficients ($\rho = -0.879, 0.286, \text{ and } -0.229$,

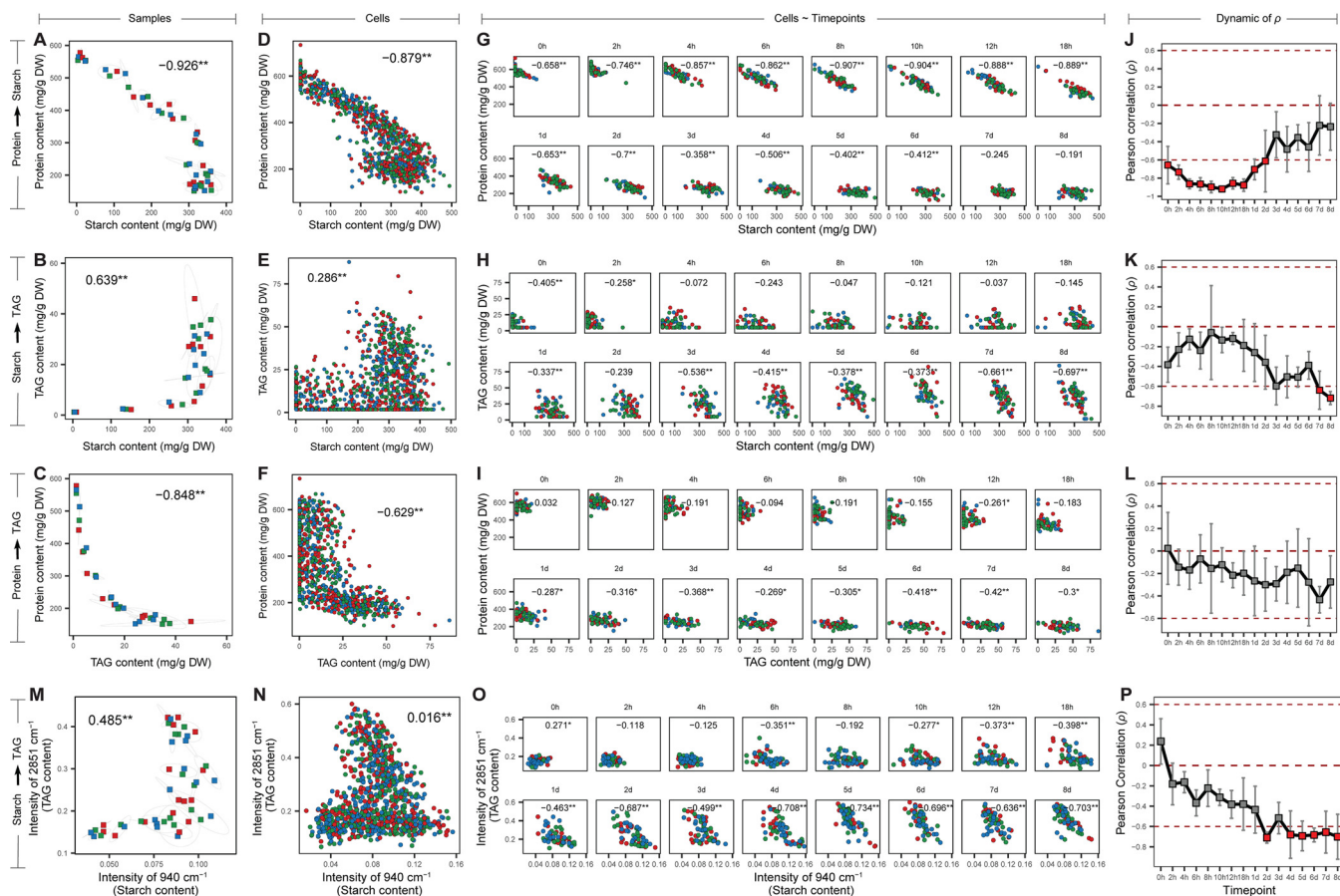


FIG 2 Correlation of starch and TAG contents of the wild-type *C. reinhardtii* CC124 at the population level and the single-cell level. (A to F) Correlation of starch and total protein contents at the population level (A) and the single-cell level (D) was shown; so was that of starch and TAG contents at the population level (B) and the single-cell level (E) and that of TAG and total protein contents at the population level (C) and the single-cell level (F). Each dot in panels A, B, and C represent one sample. Each dot in panels D, E, and F represent one individual cell. (G to L) Correlation of starch and TAG contents modeled by multiple pairs of singular Raman peaks among individual cells at each time point. Phenotypic correlation between starch (x) and protein (y) contents (G and J), between starch (x) and TAG (y) contents (H and K), and between TAG (x) and protein (y) contents (I and L) are also shown. (M to P) Phenotypic correlation between starch (x) and TAG (y) contents modeled by pair of Raman peaks of 940 cm^{-1} and 2,851 cm^{-1} . Correlation of the starch and TAG contents at the population level (M) and single-cell level (N) was compared. Phenotypic correlation between starch (x) and TAG (y) contents (O and P) at each time point were shown. In the curves of temporal dynamics, ρ is the Pearson correlation coefficients of two phenotypes among single cells (**, $P < 0.01$; *, $P < 0.05$), with value indicating the mean of triplicates and the error bar indicating the standard deviation. Absence of correlation ($\rho = 0$) or presence of strong correlation ($\rho \geq 0.6$ or $\rho \leq -0.6$) is highlighted by red horizontal lines.

respectively; Fig. 2D to F). Thus, the interphenotype correlation among pooled cells from all populations can recapitulate that among the populations.

We then probed whether such interphenotype correlation can be detected via cells from just one instance of a population, i.e., at each of the 16 time points (Fig. 2G to I). Intriguingly, for protein-starch, negative correlation (NC) is phase-specific and shows a temporal trend of gradual weakening, quite strong at each of the time points only before 2 days ($\rho \leq -0.6$, $P < 0.01$; Fig. 2G) yet absent at 7 days or 8 days (Fig. 2J). Thus, it is possible that the protein-starch conversion took place only at the early phase of N-. In fact, this prediction is supported by the routing of hydrolytic release of carbon skeletons from protein to the synthesis of starch granules during the first 2 days in *C. reinhardtii* under N-, as previously observed (22, 23).

For starch-TAG, the situation is the opposite, as the trend of NC among cells within a population intensified along time (Fig. 2H and K). Thus, in contrast to the positive correlation (PC) between starch and TAG at the population level ($\rho = 0.639$, $P < 0.01$; Fig. 2B), the within-population NC between starch and TAG was prominently present at the late phase (i.e., starting from 3 days; Fig. 2K), which predicts conversion between starch and TAG inside the cell. This single-cell-based prediction, which the population-

should attenuate in this mutant. As expected, IRCA of the time series CC4325 ramanomes revealed that the strong starch-TAG NC among single cells was no longer present (0 to 96 h under N⁻; see Fig. S2D), in contrast to CC124 (Fig. 2O and P). To validate the specificity of IRCA in revealing the starch-TAG conversion, we further collected the time series ramanomes for CC406, a wall-less strain derived from CC124 (34). For CC406, IRCA revealed strong NC ($\rho \leq -0.6$) starting at 6 h under N⁻ and remained so afterwards, suggesting starch-to-TAG conversion (see Fig. S2E). Therefore, the cell wall does not interfere with the detection of starch-TAG conversion via IRCA.

Similarly, for CC4324 (i.e., *cw15 arg7-7*), a cell wall-deficient, arginine-requiring and starch-producing strain derived from CC406, significant NC took place throughout the 96 h (especially at 24 h, 72 h, and 96 h; Fig. 3A). In contrast, for CC4333 (i.e., *sta6-1*; Fig. 3B) or CC4334 (i.e., *sta7-1*; see Fig. S3I), which both are direct CC4324 derivatives yet produce no starch due to disruption of *sta6* (ADP-glucose pyrophosphorylase) (35) and *sta7* (isoamylase) (36), respectively (34), no strong starch-TAG correlation ($\rho > -0.6$) was reported by IRCA over the full course (0 to 96 h under N⁻).

Interestingly, for CC4326 (i.e., *sta1-2*), another direct CC4324 derivative whose starch synthesis pathway is only partially disrupted (34), the temporal correlation pattern in IRCA is distinct from CC4333 and CC4334, which are fully starchless (under N⁻; Fig. 3C); instead of the latter's full ablation of starch-TAG NC throughout 96 h, CC4326 exhibited strong starch-TAG NC, yet only at the middle phase of from 12 h to 48 h, consistent with transient accumulation and conversion of a low level of starch to TAG, and then ceased the conversion upon starch depletion. Notably, under N⁺, which serves as a control condition (no TAG is accumulated in *C. reinhardtii* under abundant medium N [34]), CC4326 did not exhibit the starch-TAG NC until at the very late phase (e.g., 96 h, when the medium N was depleted by algal growth); in contrast, for CC4333 and CC4334, no starch-TAG NC was apparent throughout the 96 h under N⁺, consistent with their genotypes (i.e., complete disruption of the starch synthetic pathway; see Fig. S3 [34]).

Importantly, for both CC4565 and CC4566, both genetically complemented strains with fully restored starch synthetic capability (i.e., the *sta6* gene was reintroduced into CC4333 by overexpression via pSL18-STA6 [34]), the temporal patterns of starch-TAG NC under N⁻ are identical to those of CC4324, which is the direct progenitor of CC4333 and carries an intact starch synthetic pathway; they all exhibited significant NC throughout the 96 h (under N⁻; Fig. 3D and E). Altogether, the starch-TAG NC in IRCA that indicates starch being converted to TAG is directly correlated with starch-synthetic genotypes. Therefore, IRCA can detect and model the interconversion between metabolites from a single instance of isogenic population.

IRCNs reveal novel product-product and substrate-product links from an isogenic population. In a ramanome, the 1,581 Raman peaks in each of the many SCRS represent over one million pairwise correlations among individual cells. Such richness of information indicates an enormous number of possible between-phenotype links (and, in particular, between-metabolite conversions). For example, for the CC124 N⁻ 7-day ramanome that consists of 60 cells (which exceeds the minimal sampling depth for IRCA; see Text S1; also see Fig. S4), all pairwise among-cell correlations of peaks unveiled, in total, 60,994 strong NC "links" ($\rho \leq -0.6$, $P < 0.05$) that formulate a 1,581-node network (Fig. 4A; Materials and Methods). In such an intra-ramanome correlation network (IRCN), a node represents a Raman peak in SCRS and thus a potential phenotype (e.g., a metabolite), while an edge indicates a tentative link between two phenotypes (e.g., conversion between two metabolites).

An IRCN can reveal valuable features of the system. For example, in a 51-peak module of the CC124 N⁻ 7-day IRCN (Fig. 4B; also see Fig. S5A), besides 478 cm⁻¹ and 938 cm⁻¹ for starch and 1,658 cm⁻¹ and 2,853 cm⁻¹ for TAG, many peaks of known or unknown assignments are present, suggesting additional between-metabolite conversions. In particular, 526 cm⁻¹ (phosphatidylserine; PS) and 577 cm⁻¹ (phosphatidylinositol; PI) exhibit strong NC with 968 cm⁻¹, 1,302 cm⁻¹, and 1,263 cm⁻¹, which are all TAG markers (18) (Fig. 4B). As both PS and PI are main components of membrane

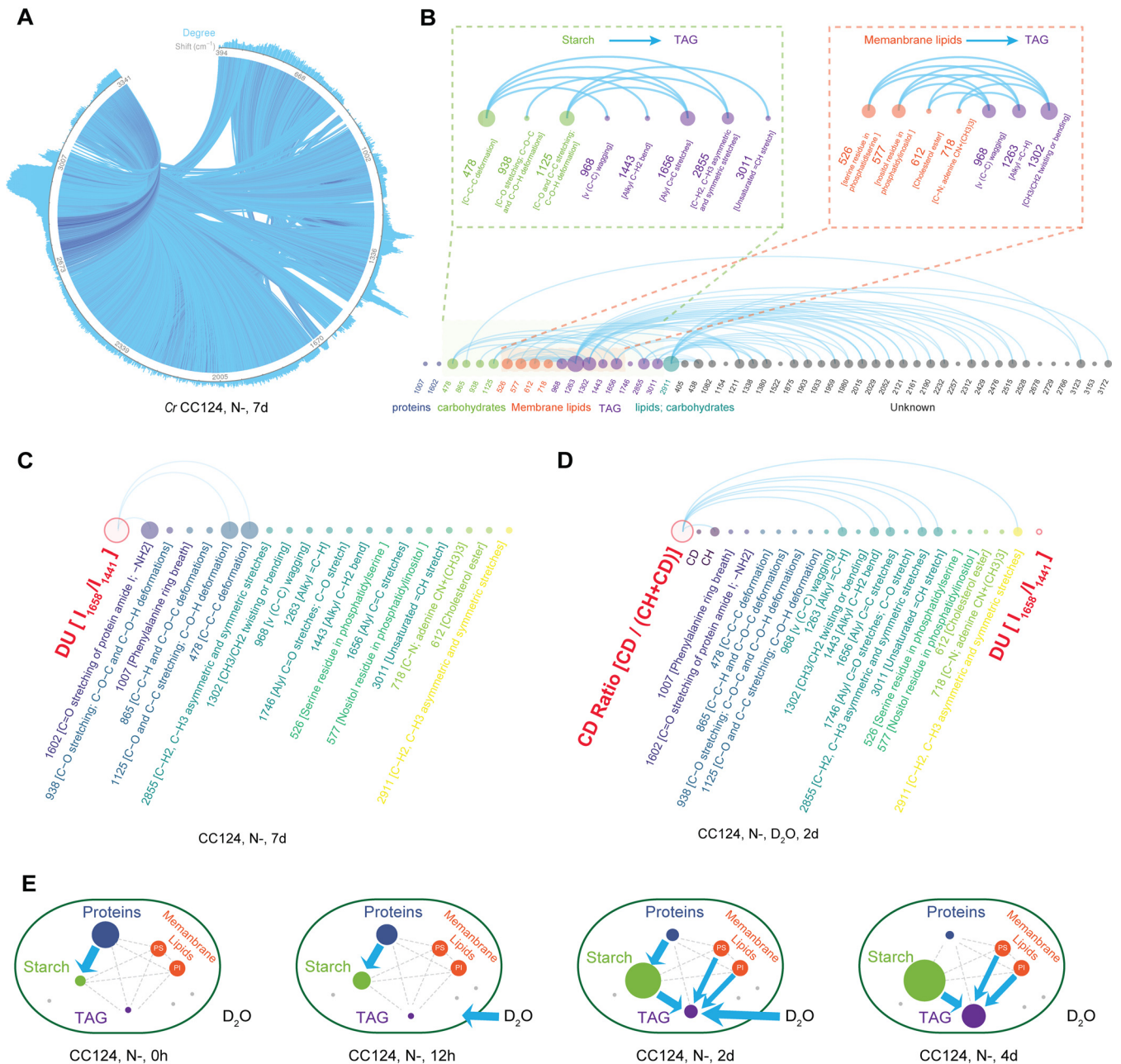


FIG 4 Global and local IRCNs of the wild-type *C. reinhardtii*. (A) The global IRCN of 1,581 Raman peaks of CC124 under N⁻ at 7 days ($\rho \leq -0.6, P < 0.05$). (B) The local IRCN with 51 Raman peaks of CC124 under N⁻ at 7 days. (C) The local IRCN (i.e., 17 characteristic Raman peaks) that includes the DU of CC124 under N⁻ at 7 days ($\rho \leq -0.6, P < 0.05$). (D) The local IRCN that further includes the CD ratio of CC124 under 25% D₂O and N⁻ at 2 days. Edges represent strong negative correlations ($\rho \leq -0.6, P < 0.05$). N⁻, nitrogen-depleted condition. (E) Key substrate-product or product-product links discovered by IRCNs. Contents of starch and TAG were quantified by 940 cm⁻¹ and 2,851 cm⁻¹, respectively. Protein content was modeled via the full SRCS. The degree of DU was quantified by the ratio between 1,658 cm⁻¹ (unsaturated C=C bonds) and 1,441 cm⁻¹ (saturated C-C bonds). The D₂O incorporation rate was quantified by the CD ratio, i.e., the ratio between the C-D bond area (2,040 to 2,300 cm⁻¹) and the area of C-D plus C-H bonds (2,040 to 2,300 cm⁻¹ and 2,800 to 3,050 cm⁻¹). Blue lines and blue arrows represent strong conversions.

lipids, these observations suggest the conversion between membrane lipids and TAG under N⁻ at 7 days (37, 38). These findings are supported by (i) the concomitant TAG accumulation and membrane lipid degradation under stress for *C. reinhardtii* (39) and (ii) the stable, modest rise of cellular FAME (fatty acid methyl ester) levels yet dramatic fall of chloroplast membrane lipid level at the early stage of N⁻ in *C. reinhardtii* (40). Notably, comparison of the 16 time series IRCNs of CC124 under N⁻ revealed a deepening trend of the PS-TAG correlation and of the PI-TAG correlation (see Fig. S5B to E),

suggesting the increasing extent of such conversions with time. Moreover, the NC between membrane lipids and TAG occurred as early as 8 h, much earlier than that between starch and TAG (Fig. 2O and P; 2 days), indicating that the membrane-lipid-TAG conversion precedes the starch-TAG conversion. On the other hand, in the N- 7-day IRCN, the starch-TAG NC is present, an indication of the starch-TAG conversion (Fig. 4B). Therefore, this local IRCN module detects multiple concomitant conversion processes that all produce TAG and reveals their dynamic features.

Notably, the IRCN can be readily expanded to include nodes representing those phenotypes that are underpinned by multiple Raman peaks. For example, the degree of unsaturation (DU), a key feature that determines the application and value of lipids, can be quantified by the ratio between the two Raman peaks of $1,658\text{ cm}^{-1}$ (unsaturated C=C bonds) and $1,441\text{ cm}^{-1}$ (saturated C-H₂ bonds) (18, 41, 42). Inclusion of DU into the aforementioned time-series IRCNs revealed a local module of 17 peaks with functional assignment, where DU is highly negatively correlated with two starch peaks (938 cm^{-1} and $1,125\text{ cm}^{-1}$) and one protein peak ($1,007\text{ cm}^{-1}$) at the late phase of N- (Fig. 4C; also see Fig. S6A and B; 2 days to 8 days). This suggests starch and proteins are converted to unsaturated lipids at the late phase of N-, consistent with the concordance between (i) degradation of proteins and starch and (ii) accumulation of unsaturated lipids (23, 28–32).

Moreover, IRCN can reveal the link between substrate intake and metabolite production of the cell. For example, cellular intake of D₂O resulted in substitution of C-H bonds in intracellular macromolecules by C-D bonds. Therefore, the CD ratio, i.e., the ratio between the C-D bond area ($2,040$ to $2,300\text{ cm}^{-1}$) and the area of C-D plus C-H bonds ($2,040$ to $2,300\text{ cm}^{-1}$ and $2,800$ to $3,050\text{ cm}^{-1}$), can measure cellular metabolic activity (43, 44). In a separate time-resolved experiment, D₂O was fed to CC124 immediately after the removal of medium nitrogen (i.e., sampled for ramanomes in triplicates at 0 h, 12 h, 1 day, 2 days, 3 days, and 4 days under N-; see Fig. S6C; Materials and Methods). IRCA revealed that (i) specifically at 2 days, a module of 17 characteristic peaks was formed, in which the CD ratio exhibited strong NC ($\rho \leq -0.6$, $P < 0.05$) with 5 TAG peaks ($1,263$, $1,443$, $1,656$, $2,855$, and $3,011\text{ cm}^{-1}$; Fig. 4D, also see Fig. S6C and D), suggesting that at 2 days, higher TAG-content cells exhibit lower D₂O-assimilating activity (and *visa versa*). Thus, at 2 days, the algal population was at a most “diverse” metabolic state, where both low-TAG, active-“drinking” cells and high-TAG, inactive-drinking cells were abundant, while before 2 days the former dominated, and after 2 days the latter dominated. (ii) Between CD ratio and DU, no strong NC ($\rho \leq -0.6$, $P < 0.05$) was observed at any of the time points (Fig. 4D, also see Fig. S6C and D), except for weak correlation at 3 days ($\rho = -0.50$, $P < 0.01$), suggesting that the DU of synthesized lipids increased with reducing *C. reinhardtii* vitality under the nitrogen-depletion stress (this finding is supported by GC-MS data [18]). (iii) In contrast to the TAG peaks, peaks of starch (the major carbon storage form of *C. reinhardtii* under N-) showed no correlation with CD ratio at any of the time points. Thus, starch synthesis appears to be mainly supported by endogenously derived H, while TAG synthesis requires exogenously supplied H. Therefore, IRCA is a new strategy to track the cellular destination of target substrate (in this case, the water).

Altogether, a choreography of interplay among water intake and major cellular products was revealed (Fig. 4E). (i) At 0 h, only the protein-starch conversion took place; (ii) at 12 h, both the protein-starch conversion and the D₂O incorporation occurred; (iii) at 2 days, besides the protein-starch conversion, TAG became the most catabolically active component, and starch, PS, PI, and D₂O all contributed to TAG synthesis; (iv) at 4 days, the three conversions of starch-TAG, PS-TAG, and PI-TAG still took place, but not the protein-starch and the D₂O-TAG conversions. Therefore, IRCA can unveil metabolite conversions from a single snapshot of an isogenic population (i.e., one ramanome), while revealing the dynamics of such conversions via a temporal series of ramanomes.

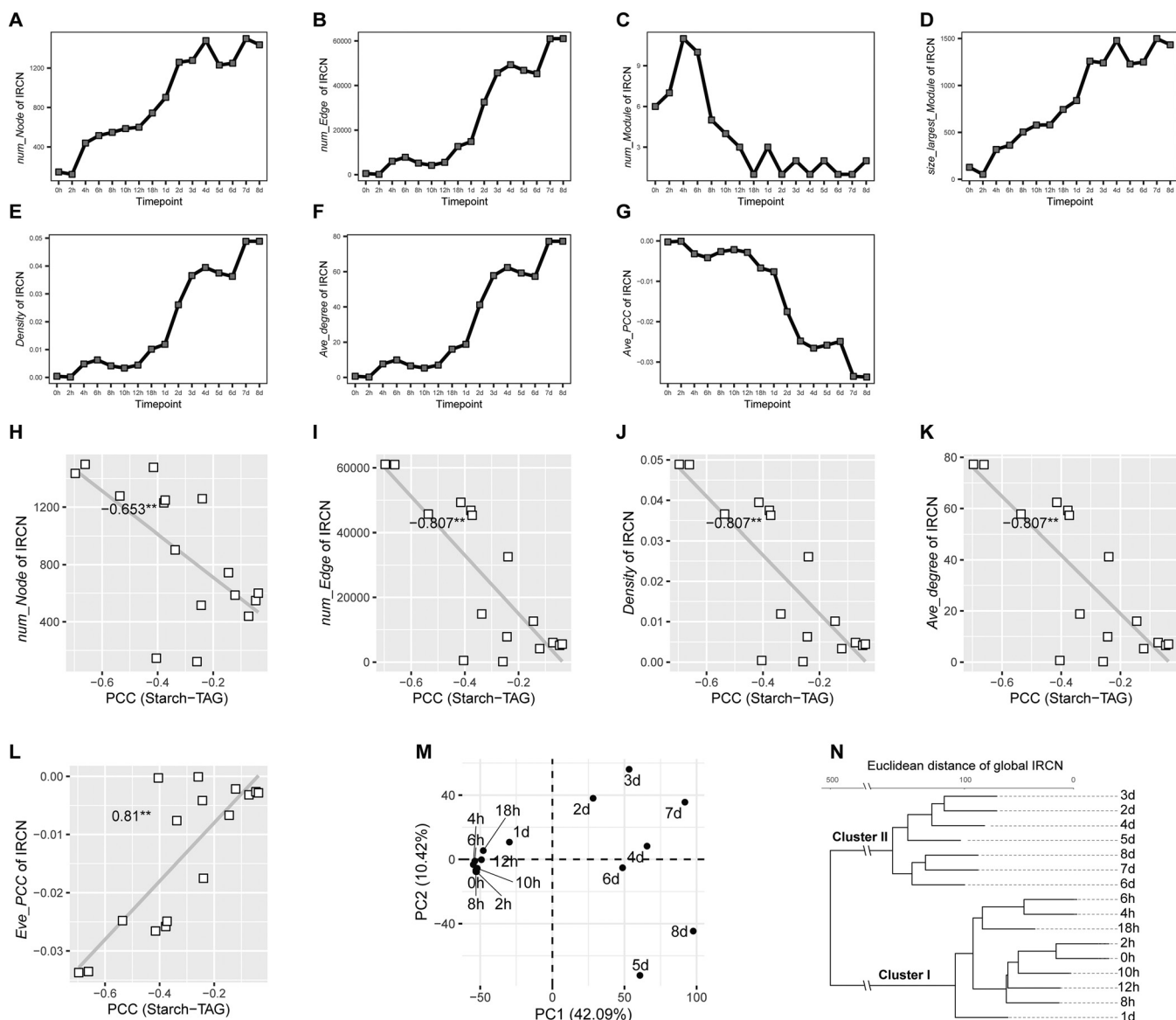


FIG 5 Intra-ramanome correlation network (IRCN) reveals global features of metabolite conversion dynamics. (A to G) Key network parameters derived from the time series IRCNs ($\rho \leq -0.6$, $P < 0.05$) of CC124 under N-, including number of nodes (A) (*num_Node*), number of edges (B) (*num_Edge*), number of modules (C) (*num_Module*; each module is a sub-IRCN not connected with any other nodes in the IRCN), size of the largest module (D) (*size_largest_Module*; number of nodes in the largest module of each IRCN), density (E) (*Density*, ratio of the number of edges divided by the number of all possible edges of the same nodes), average degree (F) (*ave_Degree*, average number of adjacent edges), and average PCC (G) (*ave_PCC*, sum of all significant strong negative correlations divided by all nodes). (H to L) Correlations between the starch-TAG conversion (i.e., PCC of starch-TAG) and key network parameters that include *num_Node* (H), *num_Edge* (I), *Density* (J), *ave_Degree* (K), and *ave_PCC* (L). (M and N) Also shown are clusters of IRCNs via PCA (M) and HCA (N) based on the correlation matrix of IRCN (strong negative correlations; $\rho \leq -0.6$, $P < 0.05$).

Global features of IRCNs reveal the degree of dynamics for metabolite conversions.

To probe the global features of IRCN, key network parameters were derived for the time series IRCNs of CC124 under N- ($\rho \leq -0.6$, $P < 0.05$; Fig. 5; heatmap of ρ shown in Fig. S7A). (i) The number of nodes (*num_Node*; Fig. 5A) increased (minimum [min] of 122 at 2 h and maximum [max] of 1,499 at 7 days), and so did the number of edges (*num_Edge*; $\rho \leq -0.6$, $P < 0.05$; Fig. 5B). (ii) The number of modules (*num_Module*; each module is a sub-IRCN that is not connected with any other nodes in the IRCN; Fig. 5C) decreased (max of 11 at 4 h and min of 1 at 18 h, 2 days, 4 days, 6 days, and 7 days). However, the size of the largest module (*size_largest_Module*; number of nodes in the largest module of each IRCN; Fig. 5D) increased greatly (min of 53 at 2 h and max of 1,499 at 7 days). (iii) Both the density (*Density*, number of edges divided by

number of all possible edges for the same nodes; Fig. 5E) and the average degree (*ave_Degree*, average number of adjacent edges; Fig. 5F), both depicting the degree of network complexity, increased. In contrast, the average Pearson correlation coefficient (*ave_PCC*; Fig. 5G) became more negative (from -0.000078 at 2 h to -0.0335 at 7 days), indicating more frequent and more active conversions between the implicated metabolites.

Across all time points, the dynamics of *num_Node*, *num_Edge*, *Density*, *ave_Degree*, and *ave_PCC* of 16 IRCNs were significantly correlated with the starch-TAG conversion (PCC of -0.653 , -0.807 , -0.807 , -0.807 , and 0.81 , respectively; $P < 0.05$; Fig. 5H to L). In addition, Raman peaks at the carbohydrate and lipid region were usually prominently found in the nodes with the highest degree (see Fig. S7B). Moreover, the $1,457\text{ cm}^{-1}$ region (the alkyl C-H₂ bend of saturated lipids) dominated at the later phases under N- (i.e., 4 days, 5 days, 7 days, and 8 days), consistent with the turnover between lipid classes (saturated switching to unsaturated). Therefore, topological features of an IRCN can reveal both metabolically active compounds and their interconversions.

Based on the matrix of ρ ($\rho \leq -0.6$, $P < 0.05$), the 16 CC124 IRCNs under N- can be classified as clusters I and II (Fig. 5M and 6N); I consists of those time points before 24 h, which are characterized by the global features of low *num_Edge* and low intensity of *ave_PCC*, and II includes those after 24 h, with global features of a sharp increase of *num_Edge* and of *ave_PCC*. Thus, 24 h is the temporal transition point (consistent with population-level measurement; see Fig. S1), when protein stopped degradation, starch stopped accumulation, and the early build-up of starch began to contribute to other cell activities, such as serving as the carbon source for lipid synthesis.

In addition, the global heatmap of $\Delta\rho$ (the difference between max ρ and min ρ in a ramanome series) reveals the dynamic change of the “hot spots” of metabolite conversion. For example, in the 16-time point CC124 N- process (see Fig. S7C), the $\Delta\rho$ between $1,441\text{ cm}^{-1}$ (alkyl C-H₂ bend, CH₂ scissoring, and CH₃ bending; i.e., saturated lipids) and $1,402\text{ cm}^{-1}$ (bending modes of methyl groups; i.e., proteins) is one of the most prominent hot spots (1.18; highlighted in Fig. S7C), with a max ρ of 0.619 at 2 h (strong positive correlation) and min ρ of -0.557 at 7 days (strong NC). This suggests the temporal switch, from the synergic degradation of lipids and protein at 2 h to the protein-to-lipid conversion at 7 d, is a key dynamic feature of this process.

Furthermore, such heatmaps of ρ can measure the global similarity of metabolite conversion modes among ramanomes from different strains (see Fig. S7D). For example, at N- 72 h, comparison among IRCNs of CC4324 (wild-type), CC4333 (starchless mutant), and CC4565 (genetically complemented strain) reveal much higher similarity with CC4324 for CC4565 than for CC4333. Specifically, CC4333 is distinct due to the absence of several regions (dotted rectangle in Fig. S7D) found in both CC4324 and CC4565, such as (i) 867 to 970 cm^{-1} (conversion of starch to lipids) and (ii) $1,441$ to $1,658\text{ cm}^{-1}$, and $1,441$ to $1,744\text{ cm}^{-1}$ (conversions of saturated lipids to unsaturated lipids; solid rectangles in Fig. S7D). These IRCN-derived phenotypes of the three strains are consistent with their genotypes (Fig. 3).

IRCN is a new “state”-specific metabolic signature of an isogenic population for diverse organisms. A ramanome is essentially a single-cell-resolution metabolic snapshot of the cellular population (16, 21). To maximally extract the metabolic features from a ramanome (Fig. 1), three signatures are proposed, metabolite profile (MP; derived via the mean SCRS of a ramanome; Fig. 6A and B), metabolite interaction (MI; derived via the correlation matrix of a ramanome consisting of significant correlations of all pairwise Raman peaks [$P < 0.05$]; Fig. 6C and D), and metabolite conversion (MC; derived via a correlation matrix of a ramanome that consists of strong NC of all pairwise Raman peaks [$\rho \leq 0.6$, $P < 0.05$]; Fig. 6E and F). To test their general applicability across organisms, MP, MI, and MC were derived for each of the 64 ramanomes from *C. reinhardtii* (wild type [WT] and the starchless mutant series), *N. oceanica*, *Saccharomyces cerevisiae*, and *Escherichia coli* (see Table S1; Materials and Methods). To facilitate interspecies comparison, the temporal ramanome data set of CC124 under N- along 16 time points

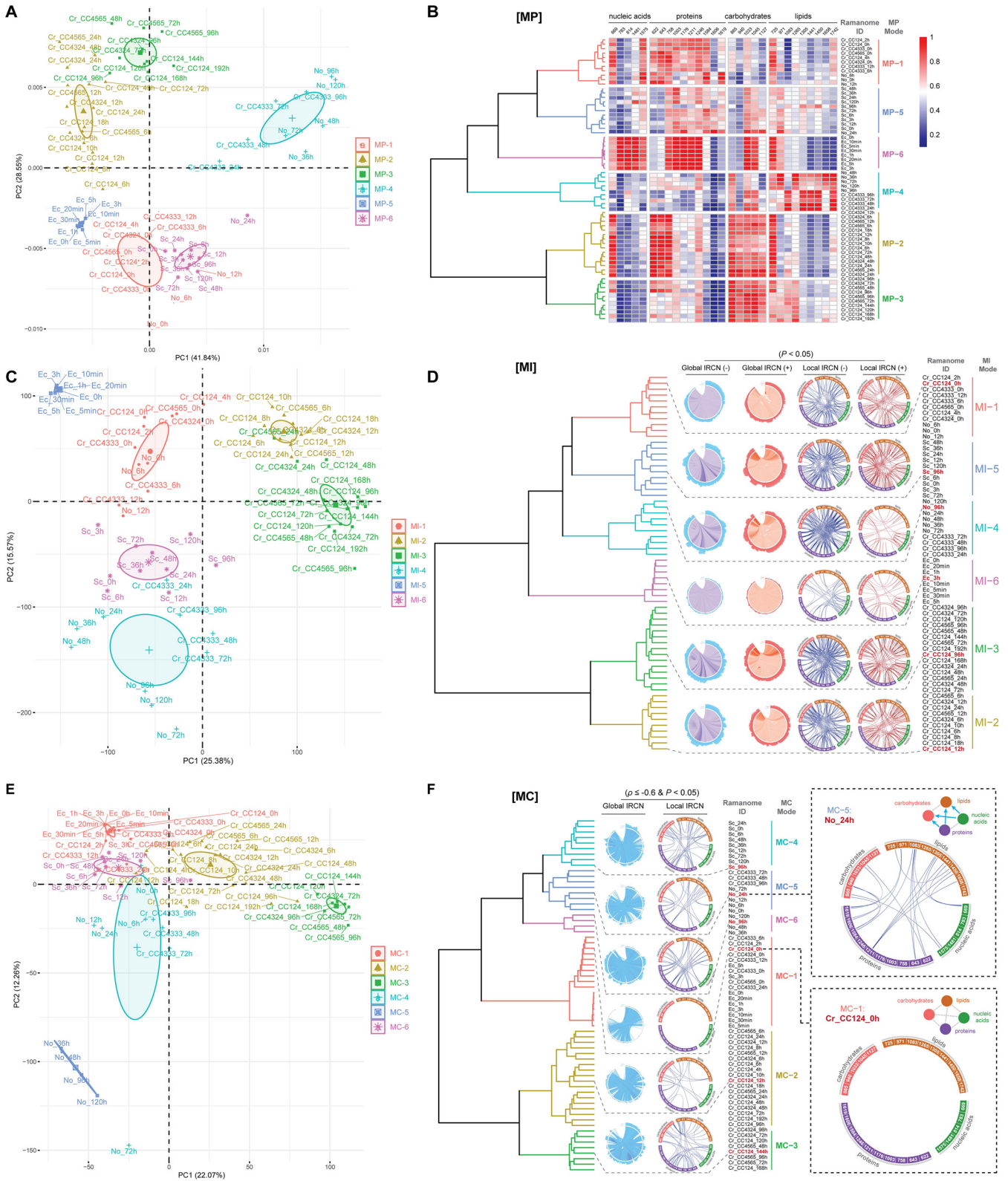


FIG 6 Clustering of ramanomes via the metabolite profiling (MP), metabolite interaction (MI), and metabolite conversion (MC) signatures. (A to F) The PCA and HCA clusters of 64 ramanomes based on MP (A and B) (mean SCRS), MI (C and D) ($P < 0.05$), and MC (E and F) ($\rho \leq -0.6$, $P < 0.05$) are shown. Details for each of the *Chlamydomonas reinhardtii* populations (WT and starchless mutant series, *C. reinhardtii* [Cr]), *Nannochloropsis oceanica* (No), *Saccharomyces cerevisiae* (Sc), and *Escherichia coli* (Ec) are provided in Table S1. Details for the Raman barcodes and local IRCNs are also provided in Table S1. Global IRCNs were derived from Raman peaks from 600 cm^{-1} to 1,800 cm^{-1} . Clusters are colored based on HCA.

was regarded as a standard “protein-starch-TAG” (PST) process, onto which the entire 64 ramanomes were “projected.”

Similarity-based clustering of MP reveals six modes (Fig. 6A and B). (i) MP-1 consists of PST 0 h, 2 h, and 4 h, plus the early phases under N⁻ of *C. reinhardtii* and *N. oceanica*, indicative of high protein content but low content of starch or TAG. (ii) MP-2 includes PST 6 h, 8 h, 10 h, 12 h, 18 h, 24 h, 48 h, and 72 h, i.e., starch-producing *C. reinhardtii* strains (CC124, CC4324, and CC4565) at the middle phases (i.e., 6 h to 48 h) under N⁻, indicating high carbohydrate content. (iii) MP-3 includes PST 96 h, 120 h, 144 h, 168 h, and 192 h, i.e., mainly starch-producing *C. reinhardtii* strains (CC124, CC4324, and CC4565) at the later phase (i.e., 48 h to 192 h) under N⁻, indicating high carbohydrate content and the start of accumulating lipids (mainly neutral lipids such as TAG). (iv) MP-4 includes both the middle and the later phases of *N. oceanica* under N⁻ (36 h, 48 h, 72 h, 96 h, and 120 h) and the later phases of starchless *C. reinhardtii* strains (CC4333) under N⁻ (24 h, 48 h, 72 h, and 96 h), corresponding to high content of neutral lipids and low levels of nucleic acids, protein, and carbohydrates. (v) MP-5 consists of *S. cerevisiae* (0 h to 120 h), featuring high protein yet low carbohydrate and lipid amounts. (vi) MP-6 consists of *E. coli* under kanamycin (0 h to 5 h), featuring larger amounts of proteins and nucleic acid.

MI forms six clusters (Fig. 6C and D). (i) MI-1 consists of 0 h, 2 h, and 4 h of PST, i.e., the very early phases under N⁻ of *C. reinhardtii* and *N. oceanica* strains, indicating that proteins began to degrade and were converted to other metabolites, such as starch. (ii) MI-2 consists of PST 6 h, 8 h, 10 h, 12 h, 18 h, and 24 h, i.e., the middle phase of starch-producing *C. reinhardtii* strains (CC124, CC4324, and CC4565) under N⁻, corresponding to high carbohydrate accumulation. (iii) MI-3 is composed of PST 48 h, 72 h, 96 h, 120 h, 144 h, 168 h, and 192 h, i.e., the later phase of starch-producing *C. reinhardtii* strains (CC124, CC4324, and CC4565) under N⁻, indicating the conversion of accumulated carbohydrates to lipids. (iv) MI-4 consists of both the middle and the later phases of *N. oceanica* under N⁻ (24 h, 36 h, 48 h, 72 h, 96 h, and 120 h) and the later phase of the starchless CC4333 under N⁻ (24 h, 48 h, 72 h, and 96 h), indicative of a state converting other metabolites such as nucleic acids, protein, and carbohydrates to lipids. (v and vi) *S. cerevisiae* (0 h to 120 h) and *E. coli* (0 h to 5 h) form MI-5 and MI-6, respectively, underscoring each species' characteristic metabolite-interacting networks.

MC also formulates six modes (Fig. 6E and F). (i) MC-1 consists of 0 h and 2 h of PST, i.e., the early phase of *C. reinhardtii* strains under N⁻ and all *E. coli* states—no or few strong pairwise NCs are found in these IRCNs. (ii) MC-2 consists of PST 4 h, 6 h, 8 h, 10 h, 12 h, 18 h, 24 h, 48 h, 72 h, and 96 h, plus the early and the middle stages of starch-producing *C. reinhardtii* mutants under N⁻, when proteins are converted into starch (see Fig. S1A). Notably, MC-2 showed more NC links between Raman peaks than MC-1, indicating more extensive metabolite conversions in MC-2. (iii) MC-3 is composed of PST 120 h, 144 h, and 168 h, i.e., the late stages of starch-producing *C. reinhardtii* strains under N⁻, when starch is converted into lipids (Fig. 2B). The NC links in IRCN of MC-3 are more abundant and stronger than those of MC-2, suggesting even more extensive and active metabolite conversions. (iv) MC-4 includes all the *S. cerevisiae* ramanome, which showed a distinct pattern of conversion among nucleic acids, proteins, carbohydrates, and lipids. (v) MC-5 mainly consists of the later phases of the starchless CC4333 and the early phase of *N. oceanica* strains under N⁻, corresponding to the metabolite-conversion pattern of lipid production. (vi) MC-6 mainly consists of the later phase of *N. oceanica* strains under N⁻, where proteins are converted into a large amount of lipids.

The MP, MI, and MC signatures, although inherently linked, can be highly distinct (see Fig. S8). Specifically, MP appears to show more species specificity, while MI and MC exhibit more state-specificity. For example, *E. coli* ramanomes are clustered with algae and yeast in MC-1; in contrast, *E. coli* ramanomes are solely clustered as MP-6. However, MC can be more sensitive in detecting cellular state change than MP and MI. For example, at 4 h under N⁻, the metabolite content of CC124 has yet to change (i.e.,

clustered with 0 h in MP-1; it did not switch to MP-2 until 6 h), yet the mode of metabolite conversion has already altered (i.e., clustered with 6 h to 96 h in MC-2).

DISCUSSION

Exploiting a fundamental and inherent property of all cellular systems, i.e., the heterogeneity in single-cell metabolism-related phenotypes (e.g., metabolite contents and substrate intake) among individual cells, here, we proposed and then biochemically and genetically validated the IRCA approach. IRCA is advantageous, as it unveils a comprehensive and landscape-like network of potential links among metabolic activities, in this case metabolite conversions or links between substrates and products, from a single instance of an isogenic population of living cells, rather than requiring a time or condition series of samples. This ability is of particular importance for those phenotyping experiments where spatiotemporal or condition-resolved sampling is a hurdle or a constraint.

Despite its need of just one snapshot of an isogenic population, an IRCN is information rich. From an IRCN, a large number of hypotheses on potential links among metabolism-related phenotypes can be generated simultaneously, based on the potentially millions of pairwise correlations from the Raman peaks that each potentially represents a phenotype or from the combinations of Raman peaks that model particular phenotypes, among the individual cells sampled for SCRS in a ramanome. The scope of such phenotypes is broad (16, 45), including but not limited to substrate intake (46), product synthesis (17, 18), and response to environmental stress (e.g., antimicrobial susceptibility [21, 44]). Based on the long and expanding list of such SCRS-derived phenotypes (16, 45), IRCNs can be constructed and then interpreted to mine the ramanome data space for new interphenotype links without *a priori* hypotheses. Moreover, since Raman microspectroscopy is label-free, noninvasive, and generally applicable to any cells, the strength of IRCA also includes high throughput, low cost, excellent scalability, and broad applicability. For example, acquisition of SCRS can be automated via flow-mode Raman cytometry or sorting (47, 48), suggesting the possibility of ultrafast, robust yet low-cost acquisition of ramanomes for IRCA. Therefore, IRCA presents a new dimension of “metabolism-related phenome” for cellular systems, which serves as a highly species- and state-specific signature of metabolic activity that captures not just the profile of Raman-sensitive metabolites (as well as other SCRS-derived phenotypes) but their dynamic links. This capacity of IRCA would enable a data-driven research strategy for profiling cellular metabolism.

On the other hand, the potential of IRCA is limited by its frequent inability to unambiguously assign Raman peaks to specific metabolites. For example, in this study, although ~1,600 spectral resonance features were detected in each SCRS, only interconversion of macromolecules such as lipids, protein, and starch can be revealed, because a large number of vibrational features were shared among metabolites and cannot be assigned uniquely to individual metabolites. Even though the list of models that link spectral resonance features to a metabolite (or other metabolism-related phenotypes) has been rapidly growing (16), only a small portion of Raman peaks in an SCRS can be assigned to a defined metabolism-related phenotype at present. In addition, biological hypotheses for many of the edges, i.e., phenotype pairs showing significant NC, in an IRCN remain difficult to interpret or confirm. Therefore, new experimental and computational methods, such as stable-isotope probed SCRS (to track the assimilation of target substrate) and multivariate curve resolution-alternating least-squares (MCR) algorithms (to deconvolute macromolecular components from overlapping Raman peaks [49, 50]) should be developed to establish new assignments or to improve their specificity for the Raman peaks, so as to enrich and expand the actual information content of an IRCN.

Despite these present limitations, as demonstrated here using a number of microalgal, fungal, and bacterial species as examples, IRCA can serve as a valuable tool to rapidly discover features of metabolic dynamics of cellular systems, such as metabolite

conversions. The capability of revealing such features from just one snapshot of an isogenic population has profound implications for designing phenotyping experiments.

MATERIALS AND METHODS

Strains and growth conditions. For *Chlamydomonas reinhardtii*, a total of nine wild-type (WT), low-starch mutant and starchless mutant strains were employed (see Table S1). Specifically, CC124, CC406, and CC4324 were WT strains. The low-starch mutant CC4325 was derived by X-ray mutagenesis from CC124. The low-starch mutant CC4326 and the starchless mutants CC4333 and CC4334 were derived from CC4324 by random integration of cassette pARG7 into the nuclear genome (34, 35, 51). The starchless mutant CC4333 is deficient in the catalytic (small) subunit of ADP-glucose pyrophosphorylase, which interrupts synthesis of the ADP-glucose, a substrate for starch biosynthesis. The CC4334 mutant contains a disrupted isoamylase gene; thus, the level of starch is severely attenuated, but it accumulates a soluble glycogen-like product. CC4565 and CC4566 were genetically complementary strains of CC4333, derived by complementation with plasmid pSL18-STA6 (34). All these mutants can be obtained from the Chlamydomonas Resource Center (<http://www.chlamycollection.org>).

The *C. reinhardtii* cells were inoculated into the TAP (Tris acetate phosphate) liquid medium with or without arginine ($100 \mu\text{g ml}^{-1}$) supplement under one-side continuous light (approximately $150 \mu\text{mol photons m}^{-2} \text{s}^{-1}$) at 25°C bubbled with air to ensure mixture and to prevent settling and were then grown to the late log phase in the N-replete TAP medium. Then they were reinoculated at 1×10^6 cells/ml in parallel into the nitrogen-replete TAP medium (N+), the nitrogen-depleted TAP medium (N-; in which NH_4Cl was omitted), or the 25% D_2O nitrogen-depleted TAP medium (25% D_2O N-), each in triplicates. Cultures at each of a series of time points were sampled in triplicate (see Table S1A).

For the *Nannochloropsis oceanica* IMET1 strain, cells were cultured in a modified f/2 liquid medium with 4 mM NO_3^- under continuous light (approximately $50 \mu\text{mol photons m}^{-2} \text{s}^{-1}$) at 25°C and then induced in nitrogen-replete (N+) or nitrogen-depleted (N-) f/2 medium. The cultures were sampled at multiple time points of 0 h, 6 h, 12 h, 24 h, 48 h, 72 h, 96 h, and 120 h. For *E. coli* DH5 α , cells were cultured in a glass tube with fresh LB medium (10g/liter NaCl, 10g/liter tryptone, 5g/liter yeast extract, $3.7 \mu\text{g/ml}$ kanamycin) at 37°C in a shaking incubator (130 rpm). Samples were collected at 0, 5, 10, 20, 30, 60, 180, and 300 min (21). For *Saccharomyces cerevisiae* Y50049, cells were cultured in a glass tube of fresh yeast extract-peptone-dextrose (YPD) medium at 30°C in a shaking incubator (200 rpm). Samples were collected at 0 h, 3 h, 6 h, 12 h, 24 h, 36 h, 48 h, 72 h, 96 h, and 120 h. All cultures and all sampling for further analysis were in triplicate.

Acquisition of single-cell Raman spectra from an isogenic population of cells. Raman spectra of individual cells were acquired using modified Raman microscopy equipped with a confocal microscope with a $\times 50$ PL magnifying dry objective (numerical aperture [NA] = 0.55, BX41; Olympus, UK) and a 532-nm Nd:YAG laser (Ventus, Laser Quantum Ltd., UK). The scattered photons were collected by a Newton electron multiplying charge coupled device (EMCCD) (Andor, UK) utilizing a $1,600 \times 200$ array of $16\text{-}\mu\text{m}$ pixels with thermoelectric cooling down to -70°C for negligible dark current. Before measurement, each sample was washed three times and resuspended in double-distilled water (ddH_2O) to remove the culture medium. For algal and yeast samples, cells were loaded into a capillary tube (50-mm length by 1-mm width by 0.1-mm height; Camlab, UK) for SCRS acquisition. The power out of the objective was 100 mW. For each SCRS, the signal acquisition time was 2 s for *C. reinhardtii* and *N. oceanica* and 3 s for yeast. For *C. reinhardtii* and *N. oceanica*, prior to Raman signal acquisition, the cell was quenched with a 532-nm laser until the signal of fluorescent and resonantly enhanced biomolecules was no longer detectable. For each individual cell, a background spectrum was generated as the average of four spectra acquired from the liquid around the cell. For *E. coli* samples, cells were loaded onto a clean CaF_2 slide and air dried before Raman measurement, and the SCRS of *E. coli* samples were collected as described (21).

Notably, a eukaryotic cell is generally larger than the laser focal spot obtained with a lens objective. In the case of a *C. reinhardtii* ($\sim 10 \mu\text{m}$ in diameter), *N. oceanica* (2 to $\sim 3 \mu\text{m}$ in diameter), or *S. cerevisiae* (2 to $\sim 3 \mu\text{m}$ in diameter) cell, which was held in the single-beam gradient force trap under the aqueous conditions, the cell was rolling in random orientation when undergoing SCRS acquisition. Therefore, the SCRS acquired represents the overall metabolic state of the cell.

Intra-ramanome correlation analysis. The raw SCRS was first preprocessed with LabSpec 5 (Horiba Scientific, France), including background subtraction and baseline correction by a polynomial algorithm (with degree of 7). Then SCRS were normalized, followed by IRCA, which used a customized computational pipeline for data analysis and result visualization.

Due to potential technical variation (e.g., change of data collector, batch, etc.), SCRS from different ramanomes may have distinct spectral ranges and resolution. Therefore, prior to computing the MP signature and the MI and MC networks, the 64 ramanomes were standardized in three steps: (i) for spectral range, only the “fingerprint area” (600 to $1,800 \text{ cm}^{-1}$) was extracted; (ii) spectral resolution was simulated to 1 cm^{-1} , via the interpolation algorithm; (iii) spectral normalization was performed by division by its area.

For each IRCN, the correlation matrix of each ramanome was constructed by calculating the Pearson correlation coefficient (PCC; ρ) of all possible pairwise combinations of Raman peaks, among all the 60 cells sampled for the ramanome. Those pairs of Raman peaks with significant correlation were considered candidates that potentially indicate links between two metabolites ($P < 0.05$), while those with strong negative correlations suggested potential conversions among two metabolites ($\rho \leq -0.6$, $P < 0.05$). The igraph package in R (<http://www.r-project.org>) was used to derive the key network

properties and visualize the IRCNs (either from specific subsets of Raman peaks or from all of the Raman peaks). To probe the global features of an IRCN, key network parameters including number of nodes (*num_Node*), number of edges (*num_Edge*), number of modules (*num_Module*; each module is a sub-IRCN not connected with any other nodes in the IRCN), size of the largest module (*size_largest_Module*), number of nodes in the largest module of each IRCN, density (*Density*, ratio of the number of edges divided by the number of all possible edges of the same nodes) and average degree (*ave_Degree*, average number of adjacent edges), and average Pearson correlation coefficient (*ave_PCC*, sum of all significant strong negative correlations divided by all nodes), were derived. For global IRCNs, all Raman peaks were used. For simplified versions of IRCN that facilitate visualization, characteristic marker Raman peaks were used (669, 783, 814, and 1,481 cm^{-1} for nucleic acids; 622, 643, 758, 1,003, 1,176, 1,211, 1,246, 1,584, 1,606, and 1,619 cm^{-1} for proteins; 865, 940, 1,033, 1,045, and 1,127 cm^{-1} for carbohydrates; and 725, 971, 1,083, 1,265, 1,305, 1,441, 1,450, 1,658, and 1,742 cm^{-1} for lipids; see Table S1B).

To characterize, compare, and cluster ramanomes, the three signatures of MP, MI, and MC were proposed. For MP, the mean SCRS of each ramanome was used. The MI network of a ramanome was generated as follows: (i) a correlation matrix was constructed by calculating the PCC of all possible pairwise Raman peaks; (ii) significant correlations ($P < 0.05$) were tabulated, while those with no significant difference were counted as 0 (no correlation). The MC network of a ramanome was generated by including only those significant, strongly negative correlations ($\rho \leq -0.6$, $P < 0.05$). To measure the similarity, pairwise Euclidean distances were calculated. Then hierarchical cluster analysis (HCA) was performed with Ward's algorithm, and six clusters were produced (52, 53). Principal-component analysis (PCA; the factextra package in R) was used to visualize the MP, MI, and MC signatures.

Data availability. The data that support the findings of this study are available from the corresponding author, Jian Xu, upon reasonable request.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

TEXT S1, DOC file, 0.3 MB.

FIG S1, TIF file, 1.4 MB.

FIG S2, TIF file, 1.9 MB.

FIG S3, TIF file, 1 MB.

FIG S4, TIF file, 0.4 MB.

FIG S5, TIF file, 0.9 MB.

FIG S6, TIF file, 1.9 MB.

FIG S7, TIF file, 2.7 MB.

FIG S8, TIF file, 1 MB.

TABLE S1, DOC file, 0.1 MB.

ACKNOWLEDGMENTS

This work was supported by Chinese Academy of Sciences (XDB29050400), the Ministry of Science and Technology of China (2018YFA0902500), the National Natural Science Foundation of China (32030003, 31900074), and the Natural Science Foundation of Shandong (ZR201807090262).

We thank Byeong-Ryool Jeong of the Ulsan National Institute of Science and Technology (UNIST) for editorial assistance.

J.X. and Y.H. introduced the IRCA concept and designed experiments. Y.H. and P.Z. performed experiments. Y.H., S.H., and Y.J. analyzed the data. J.X., and Y.H. wrote the manuscript.

We declare that we have no competing interests.

REFERENCES

- Castrillo JI, Pir P, Oliver SG. 2013. Yeast systems biology: towards a systems understanding of regulation of eukaryotic networks in complex diseases and biotechnology, p 343–365. *In* Walhout AJM, Vidal M, Dekker J (ed), *Handbook of systems biology*. Academic Press, San Diego, CA. <https://doi.org/10.1016/B978-0-12-385944-0.00018-6>.
- Wang M, Carver JJ, Phelan VV, Sanchez LM, Garg N, Peng Y, Nguyen DD, Watrous J, Kapono CA, Luzzatto-Knaan T, Porto C, Bouslimani A, Melnik AV, Meehan MJ, Liu W-T, Crüsemann M, Boudreau PD, Esquenazi E, Sandoval-Calderón M, Kersten RD, Pace LA, Quinn RA, Duncan KR, Hsu C-C, Floros DJ, Gavilan RG, Kleigrewe K, Northen T, Dutton RJ, Parrot D, Carlson EE, Aigle B, Michelsen CF, Jelsbak L, Sohlenkamp C, Pevzner P, Edlund A, McLean J, Piel J, Murphy BT, Gerwick L, Liaw C-C, Yang Y-L, Humpf H-U, Maansson M, Keyzers RA, Sims AC, Johnson AR, Sidebottom AM, Sedio BE, et al. 2016. Sharing and community curation of mass spectrometry data with global natural products social molecular networking. *Nat Biotechnol* 34:828–837. <https://doi.org/10.1038/nbt.3597>.
- Power RA, Parkhill J, de Oliveira T. 2017. Microbial genome-wide association studies: lessons from human GWAS. *Nat Rev Genet* 18:41–50. <https://doi.org/10.1038/nrg.2016.132>.
- Nicholson JK, Holmes E, Elliott P. 2008. The metabolome-wide association study: a new look at human disease risk factors. *J Proteome Res* 7: 3637–3638. <https://doi.org/10.1021/pr8005099>.

5. Go Y, Walker DI, Soltow QA, Uppal K, Wachtman LM, Strobel FH, Pennell K, Promislow DEL, Jones DP. 2015. Metabolome-wide association study of phenylalanine in plasma of common marmosets. *Amino Acids* 47:589–601. <https://doi.org/10.1007/s00726-014-1893-x>.
6. Uppal K, Soltow QA, Promislow DEL, Wachtman LM, Quyyumi AA, Jones DP. 2015. MetabNet: an R package for metabolic association analysis of high-resolution metabolomics data. *Front Bioeng Biotech* 3:87. <https://doi.org/10.3389/fbioe.2015.00087>.
7. Beck T, Shorter T, Brookes AJ. 2020. GWAS Central: a comprehensive resource for the discovery and comparison of genotype and phenotype data from genome-wide association studies. *Nucleic Acids Res* 48:933–940. <https://doi.org/10.1093/nar/gkz895>.
8. Nishino J, Ochi H, Kochi Y, Tsunoda T, Matsui S. 2018. Sample size for successful genome-wide association study of major depressive disorder. *Front Genet* 9:227. <https://doi.org/10.3389/fgene.2018.00227>.
9. Suarez-Diez M, Saccenti E. 2015. Effects of sample size and dimensionality on the performance of four algorithms for inference of association networks in metabolomics. *J Proteome Res* 14:5119–5130. <https://doi.org/10.1021/acs.jproteome.5b00344>.
10. Li J, Han D, Wang D, Ning K, Jia J, Wei L, Jing X, Huang S, Chen J, Li Y, Hu Q, Xu J. 2014. Choreography of transcriptomes and lipidomes of *Nannochloropsis* reveals the mechanisms of oil synthesis in microalgae. *Plant Cell* 26:1645–1665. <https://doi.org/10.1105/tpc.113.121418>.
11. Sanchez A, Golding I. 2013. Genetic determinants and cellular constraints in noisy gene expression. *Science* 342:1188–1193. <https://doi.org/10.1126/science.1242975>.
12. Ray LB. 2013. Cells go solo. *Science* 342:1187–1187. <https://doi.org/10.1126/science.342.6163.1187>.
13. Notingher I, Hench LL. 2006. Raman microspectroscopy: a noninvasive tool for studies of individual living cells in vitro. *Expert Rev Med Devices* 3:215–234. <https://doi.org/10.1586/17434440.3.2.215>.
14. Noothalapati Venkata Hemanth N, Shigeto S. 2012. Stable isotope-labeled Raman imaging reveals dynamic proteome localization to lipid droplets in single fission yeast cells. *Chem Biol* 19:1373–1380. <https://doi.org/10.1016/j.chembiol.2012.08.020>.
15. Huang WE, Griffiths RI, Thompson IP, Bailey MJ, Whiteley AS. 2004. Raman microscopic analysis of single microbial cells. *Anal Chem* 76:4452–4458. <https://doi.org/10.1021/ac049753k>.
16. He Y, Wang X, Ma B, Xu J. 2019. Ramanome technology platform for label-free screening and sorting of microbial cell factories at single-cell resolution. *Biotechnol Adv* 37:107388. <https://doi.org/10.1016/j.biotechadv.2019.04.010>.
17. Ji Y, He Y, Cui Y, Wang T, Wang Y, Li Y, Huang WE, Xu J. 2014. Raman spectroscopy provides a rapid, non-invasive method for quantitation of starch in live, unicellular microalgae. *Biotechnol J* 9:1512–1518. <https://doi.org/10.1002/biot.201400165>.
18. He Y, Zhang P, Huang S, Wang T, Ji Y, Xu J. 2017. Label-free, simultaneous quantification of starch, protein and triacylglycerol in single microalgal cells. *Biotechnol Biofuels* 10:275–292. <https://doi.org/10.1186/s13068-017-0967-x>.
19. Brackmann C, Norbeck J, Åkeson M, Bosch D, Larsson C, Gustafsson L, Enejder A. 2009. CARS microscopy of lipid stores in yeast: the impact of nutritional state and genetic background. *J Raman Spectrosc* 40:748–756. <https://doi.org/10.1002/jrs.2356>.
20. Collins AM, Jones HDT, Danxiang H, Qiang H, Beechem TE, Timlin JA. 2011. Carotenoid distribution in living cells of *Haematococcus pluvialis* (Chlorophyceae). *PLoS One* 6:e24302. <https://doi.org/10.1371/journal.pone.0024302>.
21. Teng L, Wang X, Xiaojun W, Honglei G, Lihui R, Tingting W, Wang Y, Yuetong J, Wei E, Huang, Xu J. 2016. Label-free, rapid and quantitative phenotyping of stress response in *E. coli* via ramanome. *Sci Rep* 6:34359. <https://doi.org/10.1038/srep34359>.
22. Msanne J, Xu D, Konda AR, Casas M, Awada T, Cahoon EB, Cerutti H. 2012. Metabolic and gene expression changes triggered by nitrogen deprivation in the photoautotrophically grown microalgae *Chlamydomonas reinhardtii* and *Coccomyxa* sp. C-169. *Phytochemistry* 75:50–59. <https://doi.org/10.1016/j.phytochem.2011.12.007>.
23. Fan J, Yan C, Andre C, Shanklin J, Schwender J, Xu C. 2012. Oil accumulation is controlled by carbon precursor supply for fatty acid synthesis in *Chlamydomonas reinhardtii*. *Plant Cell Physiol* 53:1380–1390. <https://doi.org/10.1093/ppcp/pcs082>.
24. Li Y, Han D, Hu G, Dauvillee D, Sommerfeld M, Ball S, Hu Q. 2010. *Chlamydomonas* starchless mutant defective in ADP-glucose pyrophosphorylase hyper-accumulates triacylglycerol. *Metab Eng* 12:387–391. <https://doi.org/10.1016/j.ymben.2010.02.002>.
25. Goodenough U, Blaby I, Casero D, Gallaher SD, Goodson C, Johnson S, Lee J, Merchant SS, Pellegrini M, Roth R, Rusch J, Singh M, Umen JG, Weiss TL, Wulan T. 2014. The path to triacylglyceride obesity in the *sta6* strain of *Chlamydomonas reinhardtii*. *Eukaryot Cell* 13:591–613. <https://doi.org/10.1128/EC.00013-14>.
26. Work VH, Radakovits R, Jinkerson RE, Meuser JE, Elliott LG, Vinyard DJ, Laurens LML, Dismukes GC, Posewitz MC. 2010. Increased lipid accumulation in the *Chlamydomonas reinhardtii* *sta7-10* starchless isoamylase mutant and increased carbohydrate synthesis in complemented strains. *Eukaryot Cell* 9:1251–1261. <https://doi.org/10.1128/EC.00075-10>.
27. Johnson X, Alric J. 2013. Central carbon metabolism and electron transport in *Chlamydomonas reinhardtii*: metabolic constraints for carbon partitioning between oil and starch. *Eukaryot Cell* 12:776–793. <https://doi.org/10.1128/EC.00318-12>.
28. Fan J, Ning K, Zeng X, Luo Y, Wang D, Hu J, Li J, Xu H, Huang J, Wan M, Wang W, Zhang D, Shen G, Run C, Liao J, Fang L, Huang S, Jing X, Su X, Wang A, Bai L, Hu Z, Xu J, Li Y. 2015. Genomic foundation of starch-to-lipid switch in oleaginous *Chlorella* spp. *Plant Physiol* 169:2444–2461. <https://doi.org/10.1104/pp.15.01174>.
29. Vigeolas H, Möhlmann T, Martini N, Neuhaus HE, Geigenberger P. 2004. Embryo-specific reduction of ADP-Glc pyrophosphorylase leads to an inhibition of starch synthesis and a delay in oil accumulation in developing seeds of oilseed rape. *Plant Physiol* 136:2676–2686. <https://doi.org/10.1104/pp.104.046854>.
30. Andriotis VME, Pike MJ, Kular B, Rawsthorne S, Smith AM. 2010. Starch turnover in developing oilseed embryos. *New Phytol* 187:791–804. <https://doi.org/10.1111/j.1469-8137.2010.03311.x>.
31. Heredia-Arroyo T, Wei W, Hu B. 2010. Oil accumulation via heterotrophic/mixotrophic *Chlorella* protothecoides. *Appl Biochem Biotechnol* 162:1978–1995. <https://doi.org/10.1007/s12010-010-8974-4>.
32. Wei X, Liu L, Chao W, Chen Y, Wu Q. 2010. ¹³C-tracer and gas chromatography-mass spectrometry analyses reveal metabolic flux distribution in the oleaginous microalga *Chlorella* protothecoides. *Plant Physiol* 154:1001–1011. <https://doi.org/10.1104/pp.110.158956>.
33. Ball S, Marianne T, Dirick L, Fresnoy M, Delrue B, Decq A. 1991. A *Chlamydomonas reinhardtii* low-starch mutant is defective for 3-phosphoglycerate activation and orthophosphate inhibition of ADP-glucose pyrophosphorylase. *Planta* 185:17–26. <https://doi.org/10.1007/BF00194509>.
34. Siaux M, Cuiné S, Cagnon C, Fessler B, Nguyen M, Carrier P, Beyly A, Beisson F, Triantaphylidès C, Li-Beisson Y, Peltier G. 2011. Oil accumulation in the model green alga *Chlamydomonas reinhardtii*: characterization, variability between common laboratory strains and relationship with starch reserves. *BMC Biotechnol* 11:7. <https://doi.org/10.1186/1472-6750-11-7>.
35. Zabawinski C, Van den Koornhuysen N, D'Hulst C, Schlichting R, Giersch C, Delrue B, Lacroix J, Preiss J, Ball S. 2001. Starchless mutants of *Chlamydomonas reinhardtii* lack the small subunit of a heterotetrameric ADP-glucose pyrophosphorylase. *J Bacteriol* 183:1069–1077. <https://doi.org/10.1128/JB.183.3.1069-1077.2001>.
36. Mouille G, Maddelein M, Libessart N, Talaga P, Decq A, Delrue B, Ball S. 1996. Preamylopectin processing: a mandatory step for starch biosynthesis in plants. *Plant Cell* 8:1353–1366. <https://doi.org/10.1105/tpc.8.8.1353>.
37. Krafft C, Neudert L, Simat T, Salzer R. 2005. Near infrared Raman spectra of human brain lipids. *Spectrochim Acta A Mol Biomol Spectrosc* 61:1529–1535. <https://doi.org/10.1016/j.saa.2004.11.017>.
38. Movasaghi Z, Rehman S, Rehman IU. 2007. Raman spectroscopy of biological tissues. *Appl Spectrosc Rev* 42:493–541. <https://doi.org/10.1080/05704920701551530>.
39. Yoon K, Han D, Li Y, Sommerfeld M, Hu Q. 2012. Phospholipid:diacylglycerol acyltransferase is a multifunctional enzyme involved in membrane lipid turnover and degradation while synthesizing triacylglycerol in the unicellular green microalga *Chlamydomonas reinhardtii*. *Plant Cell* 24:3708–3724. <https://doi.org/10.1105/tpc.112.100701>.
40. Juergens MT, Disbrow B, Shachar-Hill Y. 2016. The relationship of triacylglycerol and starch accumulation to carbon and energy flows during nutrient deprivation in *Chlamydomonas*. *Plant Physiol* 171:2445–2457. <https://doi.org/10.1104/pp.16.00761>.
41. Ota S, Alexandr J, Zdeněk P, Pavel Z, Ladislav N, Jan T, Petr K, Martin T. 2010. Raman microspectroscopy of individual algal cells: sensing unsaturation of storage lipids in vivo. *Sensors* 10:8635–8651. <https://doi.org/10.3390/s100908635>.

42. Wu H, Volponi JV, Oliver AE, Parikh AN, Simmons BA, Seema S. 2011. In vivo lipidomics using single-cell Raman spectroscopy. *Proc Natl Acad Sci U S A* 108:3809–3814. <https://doi.org/10.1073/pnas.1009043108>.
43. Berry D, Mader E, Lee TK, Woebken D, Wang Y, Zhu D, Palatinszky M, Schintlmeister A, Schmid MC, Hanson BT. 2014. Tracking heavy water (D2O) incorporation for identifying and sorting active microbial cells. *P Natl Acad Sci U S A* 112:194–203. <https://doi.org/10.1073/pnas.1420406112>.
44. Tao Y, Wang Y, Huang S, Zhu P, Huang WE, Ling JQ, Xu J. 2017. Metabolic-activity based assessment of antimicrobial effects by D2O-labeled single-cell Raman microspectroscopy. *Anal Chem* 89:4108–4115. <https://doi.org/10.1021/acs.analchem.6b05051>.
45. Wang Y, Song Y, Tao Y, Muhamadali H, Goodacre R, Zhou N, Preston GM, Xu J, Huang WE. 2016. Reverse and multiple stable isotope probing to study bacterial metabolism and interactions at the single cell level. *Anal Chem* 88:9443–9450. <https://doi.org/10.1021/acs.analchem.6b01602>.
46. Hatzenpichler R, Krukenberg V, Spietz R, Jay Z. 2020. Next-generation physiology approaches to study microbiome function at single cell level. *Nat Rev Microbiol* 18:241–256. <https://doi.org/10.1038/s41579-020-0323-1>.
47. Hiramatsu K, Ideguchi T, Yonamine Y, Lee S, Luo Y, Hashimoto K, Ito T, Hase M, Park JW, Kasai Y, Sakuma S, Hayakawa T, Arai F, Hoshino Y, Goda K. 2019. High-throughput label-free molecular fingerprinting flow cytometry. *Sci Adv* 5:eaau0241. <https://doi.org/10.1126/sciadv.aau0241>.
48. Wang X, Xin Y, Ren L, Sun Z, Zhu P, Ji Y, Li C, Xu J, Ma B. 2020. Positive dielectrophoresis-based Raman-activated droplet sorting for culture-free and label-free screening of enzyme function in vivo. *Sci Adv* 6:eabb3521. <https://doi.org/10.1126/sciadv.abb3521>.
49. Ando M, Hamaguchi H. 2014. Molecular component distribution imaging of living cells by multivariate curve resolution analysis of space-resolved Raman spectra. *J Biomed Opt* 19:e011016. <https://doi.org/10.1117/1.JBO.19.1.011016>.
50. Noothalapati H, Iwasaki K, Yamamoto T. 2017. Biological and medical applications of multivariate curve resolution assisted Raman spectroscopy. *Anal Sci* 33:15–22. <https://doi.org/10.2116/analsci.33.15>.
51. Li Y, Han D, Hu G, Sommerfeld M, Hu Q. 2010. Inhibition of starch synthesis results in overproduction of lipids in *Chlamydomonas reinhardtii*. *Biotechnol Bioeng* 107:258–268. <https://doi.org/10.1002/bit.22807>.
52. Lu W, Chen X, Wang L, Li H, Fu YV. 2020. Combination of an artificial intelligence approach and laser tweezers Raman spectroscopy for microbial identification. *Anal Chem* 92:6288–6296. <https://doi.org/10.1021/acs.analchem.9b04946>.
53. Choo-Smith LP, Maquelin K, van Vreeswijk T, Bruining HA, Puppels GJ, Ngo Thi NA, Kirschner C, Naumann D, Ami D, Villa AM, Orsini F, Doglia SM, Lamfarraj H, Sockalingum GD, Manfait M, Allouch P, Endtz HP. 2001. Investigating microbial (micro)colony heterogeneity by vibrational spectroscopy. *Appl Environ Microbiol* 67:1461–1469. <https://doi.org/10.1128/AEM.67.4.1461-1469.2001>.