

# *IN SILICO* SINGLE-CELL METABOLISM ANALYSIS UNRAVELS A NEW TRANSITION STAGE OF CD8 T CELLS 4 DAYS POST-INFECTION

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## **ABSTRACT**

CD8 T cell proper differentiation during antiviral responses relies on metabolic adaptations. Herein, we investigated global metabolic activity in single CD8 T cells along an *in vivo* response by estimating metabolic fluxes from single-cell RNA-sequencing data. The approach was validated by the observation of metabolic variations known from experimental studies on global cell populations, while adding temporally detailed information and unravelling yet undescribed sections of CD8 T cell metabolism that are affected by cellular differentiation. Furthermore, inter-cellular variability in gene expression level, highlighted by single cell data, and heterogeneity of metabolic activity 4 days post-infection, revealed a new transition stage accompanied by a metabolic switch in activated cells differentiating into full-blown effectors.

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## INTRODUCTION

CD8 T lymphocytes are critical cytotoxic effector cells that protect against viral infections by eradicating virus-infected cells. The antiviral response typically recruits rare antigen (Ag)-specific naive CD8 T cells, resulting in their tremendous proliferation and differentiation into potent cytotoxic cells that eradicate infected cells. After clearing of viral Ags, the majority of effectors undergoes apoptosis leaving a small population of quiescent memory cells, which will ensure rapid and most effective secondary responses to subsequent infections with the same pathogen (Murali-Krishna et al., 1998; Kaech and Cui, 2012).

This differentiation process, which produces immediate effector and long-term protector lymphocyte populations, is accompanied by metabolic reprogramming at different stages in order to support specific bioenergetic requirements of differentiating cells (Klein Geltink et al., 2018; Gupta et al., 2020; Møller et al., 2022). For instance, glucose consumption by CD8 T lymphocytes relies on its cytoplasmic degradation to pyruvate that can be further catabolized to lactate, a process known as glycolysis. Alternatively, pyruvate can enter mitochondria, where its conversion into acetyl-coenzymeA (acetyl-coA) fuels the TriCarboxylic Acid (TCA), which in turn activates the mitochondrial membrane Electron Transport Chain (ETC) for Oxidative phosphorylation (Oxphos) of Adenosine-DiPhosphate (ADP) to Adenosine-TriPhosphate (ATP). CD8 T cell can also produce energy from fatty acids (FA)-oxidation (FAO) in mitochondria, which produces acetyl-coA for the TCA. The metabolism of quiescent naive CD8 T cells mostly relies on basic glycolysis and FAO (Gupta et al., 2020). Upon activation, CD8 T cells enhance glucose consumption (Menk et al., 2018) and switch to aerobic glycolysis (van der Windt and Pearce, 2012; Klein Geltink et al., 2018; Salmond, 2018; Jung et al., 2019; Gupta et al., 2020; Møller et al., 2022). This metabolic switch is mandatory for the proper differentiation of naive cells into effectors (Pearce et al., 2013; Pollizzi et al., 2015). Later in the response, when effectors differentiate to memory cells, they undergo a new metabolic switch back to FAO (Araki et al., 2009; Pearce et al., 2009; van der Windt and Pearce, 2012), which is again mandatory for proper memory CD8 T cell generation (Pollizzi et al., 2015; Bevilacqua et al., 2022; Møller et al., 2022).

Besides this well described reprogramming of energy production during CD8 T cell responses to viral infections, Ag encounter also promotes cholesterol biosynthesis (Chen et al., 1975; Kidani et al., 2013) and triggers an increase in glutamine uptake and glutaminolysis (Carr et al., 2010; Wang et al., 2011), which fuels FA synthesis through  $\alpha$ -KetoGlutarate ( $\alpha$ -KG)-dependent citrate production (Gupta et al., 2020; Pearce et al., 2013). T cell activation also redirects glyceraldehyde-3P (G3P) downstream glucose degradation towards the production of

5-phosphoribose-2P (PRPP) that fuels the pentose-phosphate pathway (PPP) to meet nucleic acid and aromatic amino acid biosynthesis demands (Wang et al., 2011). Globally, CD8 T cell activation, proliferation and differentiation are coupled to metabolic reprogramming that support trade-offs between energy demand and biomolecule synthesis and participate in cell differentiation. Both cell differentiation and metabolic switches are intertwined and co-regulated epigenetically (Almeida et al., 2016; Jung et al., 2019; Yerinde et al., 2019; Gupta et al., 2020; Møller et al., 2022).

These metabolic adaptations to viral challenge have been studied on global populations of CD8 T cell responders. However, effector and memory CD8 T cell populations are heterogeneous (Appay et al., 2002; Kaech and Cui, 2012; Mittrücker et al., 2014) and different subsets shall undergo and require specific metabolic programs (Geiger et al., 2016; Gupta et al., 2019). Thus, an analysis of the metabolism of CD8 T cell responding to a viral infection at the single-cell level would be much more biologically relevant (Arsenio et al., 2014; Ahl et al., 2020; Fernández-García et al., 2022). However, single-cell metabolomic techniques still suffer from relatively low throughput and sensitivity (Duncan et al., 2019; Sengupta et al., 2019). Thus, to generate a global description of the metabolism of CD8 T cell responding to a viral infection at the single-cell level, we made use of the recently described single-cell Flux Estimation Analysis (scFEA) algorithm (Alghamdi et al., 2021) that estimates metabolite fluxes in cells from single-cell RNA-sequencing (scRNA-seq) expression data. Our results evidenced metabolic switches previously described, thus validating the approach, and revealed new metabolic perturbations that may deserve further detailed experimental validation.

Furthermore, this single-cell level analysis revealed time-dependent variation in inter-cellular heterogeneity. Indeed, we show that, as in numerous other cell differentiation systems (Papili Gao et al., 2020), metabolic genes are subjected to a transient rise in gene expression variability allowing cells to explore the gene expression space, before selection of the most appropriate gene regulatory network (GRN) state resulting in a more homogeneous gene expression pattern in the emerging differentiated populations (Guillemin and Stumpf, 2021). Interestingly, two distinct surges in metabolic gene expression variability were observed, immediately after activation and around 4 days post-infection (dpi). The genes concerned by these perturbations are involved in largely non-overlapping pans of cell metabolism, suggesting a yet unknown metabolic switch at 4 dpi, when activated cells fully commit to effector differentiation. This switch may be coupled to a selection process, as the scFEA analysis revealed that many cells at 4 dpi could not sustain the metabolic changes triggered by cell activation. Finally, it occurs at a transition stage globally affecting the GRN, beyond metabolic regulation.

# RESULTS

## Expression data

We obtained expression data from a scRNA-seq study of murine CD8 T cells responding to an acute viral infection (Kurd et al., 2020). Therein, the authors transferred P14 transgenic CD8 T cells that recognize an LCMV epitope to histocompatible hosts, which were acutely immunized with the virus the day after. Single responding CD8 T cells from the spleens of immunized hosts were then sorted at different days post-infection (dpi) and analyzed by scRNA-seq. We performed quality control of the data downloaded from Gene Expression Omnibus (GSE131847) and selected cells and genes (Methods) to generate a final UMI count table of 14,666 genes in 42,025 single cells (Figure 1A). Dimension reduction and representation on UMAP revealed a correct temporal arrangement of the cells (Figure 1B), although D4 and D32 cells split in two regions of the projection.

## Global analysis of metabolic changes in responding CD8 T cells

In order to have access to an unbiased global analysis of the metabolism in CD8 T cells responding to an *in vivo* viral infection at the single-cell level, we used the scFEA algorithm that can estimate metabolic fluxes from scRNA-seq expression data (Alghamdi et al., 2021). The authors have subdivided all the metabolic pathways of human and murine cells into 171 flux-independent modules of enzymatic reactions. These modules are then reconstructed as a factor graph based on the network topology and gene expression status. Finally, all cell fluxomes are estimated thanks to a multilayer neural network model to capture the nonlinear dependency of metabolic fluxes on the enzymatic gene expressions.

## Fluxome analysis

We selected 530 metabolic genes (Methods) for scFEA analysis. As expected, their level of expression varies during the primary T cell response (SupplementaryFigure 1). We submitted UMI counts for these 530 metabolic genes in the 42,025 cells to scFEA and the algorithm returned flux values in all cells for 168 of the designed modules. We selected modules with relevant flux values (Methods) and did not consider the modules corresponding to simple metabolite efflux from cells (Super-Module 12 in Alghamdi et al. (2021)). Modules with a substantial variation of flux along the differentiation kinetics were further selected (Methods), resulting in a list of 34 metabolic modules of interest (SupplementaryTable 2, SupplementaryFigure 4). Those belong to 13 of the 22 Super metabolic Modules (SM) defined by Alghamdi et al. (2021), including SM1 corresponding to ‘glycolysis and TCA cycle’, which

are crucial for CD8 T cell responses. Furthermore, spermine, beta-alanine, leucine+valine+isoleucine and spermine metabolisms; the pentose phosphate pathway; the urea cycle and the synthesis of purines, pyrimidines, N- and O-linked glycans, as well as steroid hormones, all displayed significant and varying fluxes (SupplementaryTable 2).

Flux variations all occur between D0 and D10 of the response with a surge in variation at D5, all fluxes returning to their initial values by D10 and remaining stable up to the memory phase of the response at D90 (SupplementaryFigure 4). The flux variations display 3 typical patterns: fluxes either increase (Figure 2A) or decrease (Figure 2B) between D0 and D10, or show a biphasic evolution with a transient decrease between D0 and D4, followed by an increase above initial values from 5 to 10 dpi (Figure 2C). Whatever the direction of the flux variations, cells at 4 dpi are heterogeneous with a majority of cells displaying a flux corresponding to the variation between 3 and 5 dpi, but a minority of cells with a flux similar to initial values, as if they were not able to sustain the initial metabolic changes.

The projection of the fluctuating flux modules on the murine metabolic map from KEGG reveals a regionalization of the selected modules by Super-Module (Figure 3) and complex variations of fluxes during the differentiation of cells. For instance, in the 'Pyrimidine synthesis' SM21, the flux through M-155 that corresponds to the reactions Uridine-TriPhosphate (UTP)  $\rightarrow$  Cytidine-TriPhosphate (CTP)  $\rightarrow$  Cytidine-DiPhosphate (CDP) is upregulated between D0 and D10, while the flux through M-153 that corresponds to the overlapping reaction Uridine-MonoPhosphate (UMP)  $\rightarrow$  UTP  $\rightarrow$  CTP  $\rightarrow$  CDP is initially downregulated between D0 and D4 and then upregulated (SupplementaryFigure 5A). Furthermore, the fluxes through M-157 (CDP  $\rightarrow$  deoxyCDP (dCDP)), M-158 (dCDP  $\rightarrow$  deoxycytidine-MonoPhosphate (dCMP)) and M-171 (dCMP  $\rightarrow$  deoxycytidine) are up regulated, while the flux through M-161 (dCDP  $\rightarrow$  deoxycytidine-TriPhosphate (dCTP)) is downregulated. Such complex flux interactions are difficult to interpret in terms of metabolite concentrations, such as for CDP or dCDP, and were evidenced in other Super-Modules: SM17 (SupplementaryFigure 5B) and SM16 (SupplementaryFigure 5C).

### **Metabolic stress analysis**

Thus, in order to relate flux variations in modules and metabolite accumulation or deprivation from cells, we again used scFEA that calculates metabolite concentrations from the estimated fluxes. scFEA returned values for 70 metabolites. We selected (Methods) the top19 variable metabolites for analysis (SupplementaryTable 3). The concentration of metabolites varied between D0 and D14 of the response (Figure 4) with a peak at 5 dpi. Cells were transiently deprived of Glucose-6-Phosphate (G6P), pyruvate, acetyl-coA and glycan backbones

(GlcNAc)<sub>4</sub> (Man)<sub>3</sub> (Asn)<sub>1</sub> and (Gal)<sub>1</sub> (GlcNAc)<sub>1</sub> (Man)<sub>1</sub> (Ser/Thr)<sub>1</sub>; and transiently accumulated leucine, isoleucine, lactate and deoxyadenosine. For these metabolites, a large fraction of cells at 4 dpi did not sustain the accumulation or loss, as observed for fluxes (Figure 2). Other metabolite concentrations were biphasic with G3P, glutathione, E,E-Farnesyl-PP, UMP and dolichyl-P-D-mannose being accumulated up to 4 dpi and then deprived from cells. Conversely, after an initial loss from D0 to D4, cells accumulated glutamate, cholesterol and 5-Phosphoribose-2P (PRPP) between D5 and D14. Finally, CDP concentration displayed oscillations rising at D3, then decreasing below initial values with a nadir at 5 dpi, followed by a new maximum at D6, while dCDP concentration oscillated in mirror (Figure 4).

Interestingly, when the top19 variable metabolites were localized on the metabolic map, they all matched a varying flux module (Figure 3). Indeed, Glutamate, PRPP and cholesterol, which are the products of M-25, M-33, M-167 modules with first decreasing and then up-regulated fluxes, were first depleted from cells before accumulating, while Glutathione, G3P and (E,E)-Farnesyl-PP, the substrates of these same modules were first accumulated before loss. Also, leucine and isoleucine substrates of M-53 and M-55/M-56 modules with decreasing fluxes and deoxyadenosine product of M-140 with an increasing flux accumulated, while the increase of M-6 flux from pyruvate to lactate resulted in an accumulation of lactate and pyruvate depletion. Furthermore, the analysis of metabolite stress helped interpret the complex interactions of varying fluxes. For instance, the complex interactions of fluxes in all 6 modules of the ‘pyrimidine synthesis’ SM21 resulted in opposite oscillations in CDP and dCDP concentrations, while fluctuations in M-153 flux resulted in an accumulation of UMP prior to depletion (SupplementaryFigure 7A). Similarly, the rise in M-126 flux counteracted fluctuations in M-125 flux and the increase in M-122 competed the decrease and fluctuations of M-119 and M-121, so that cells were depleted from (Gal)<sub>1</sub> (GlcNAc)<sub>1</sub> (Man)<sub>1</sub> (Ser/Thr)<sub>1</sub> and (GlcNAc)<sub>4</sub> (Man)<sub>3</sub> (Asn)<sub>1</sub> glycans (SupplementaryFigure 7B, C). This is in accordance with the stronger values of fluxes in M-126 and M-122, as compared to fluxes in M-125 and M-119 and M-121, respectively (SupplementaryTable 2).

In a whole, scFEA analysis led to a very detailed global and temporal single-cell description of variations in metabolic fluxes and metabolite concentrations in CD8 T cells during a viral infection.



## Stochasticity of metabolic gene expression

Single-cell resolution of metabolic fluxome analysis showed a large degree of inter-cellular heterogeneity, revealing a cell subset at 4 dpi (Figure 2, SupplementaryFigure 4) and highlighting the importance of this degree of granularity to characterize CD8 T cell responses. We next questioned whether inter-cellular heterogeneity in metabolic fluxes stems from a certain degree of stochasticity in the differentiation process. Indeed, in most cell differentiation contexts, a rise in the stochasticity of gene expression has been observed just prior to transition or branching points in the differentiation process (Richard et al., 2016; Stumpf et al., 2017; Papili Gao et al., 2020; Dussiau et al., 2022; Parmentier et al., 2022; Toh et al., 2022). Such an increase in gene expression variability allows cells submitted to environmental perturbations to explore the gene expression space before selection of the most appropriate GRN state in the resulting differentiated population (Huang et al., 2005; Huang, 2009). This phenomenon has not yet been described for mature lymphocytes differentiating in response to an antigenic challenge. As shown in SupplementaryFigure 1, the level of metabolic gene expression varies during CD8 T cell differentiation. However, during hematopoiesis, if many genes show fluctuations in expression level, it is the ones that show the highest fluctuations in cell-to-cell variability over the course of a differentiation trajectory that are pathway-specific (Dussiau et al., 2022). Since metabolic regulation is crucial for the proper differentiation of CD8 T cells in response to a viral challenge, we thus assessed the inter-cellular variability in metabolic gene expression during this differentiation process.

For this, we estimated the degree of inter-cellular variability in the expression of the 530 metabolic genes selected for flux estimations by their level of entropy (Dussiau et al., 2022; Gandrillon et al., 2021) at each dpi (Methods). As shown in Figure 5A, many metabolic genes show strong variations in expression variability, mostly during the first 10 days after infection. Thus, although metabolic gene expression is strictly regulated during CD8 T cell differentiation (Gupta et al., 2019; Møller et al., 2022), surges in inter-cellular variability indicate the process is submitted to variation in stochasticity. There is a weak correlation ( $r=0.68$ ) between the variations in gene expression and entropy (SupplementaryFigure 8), as previously seen in other cell differentiation models (Dussiau et al., 2022). In order to highlight kinetic patterns of metabolic gene entropies, we clustered entropy profiles shown in Figure 5A, with functional PCA followed by kmeans clustering (Methods). We identified two groups of patterns in metabolic gene expression variabilities (Figure 5B, C): 86 genes (SupplementaryTable 4) show a strong, transient and immediate surge in entropy after activation, hereafter called profile 1,

while 444 genes show a weaker and later increase in entropy, beginning at 4 dpi, with a peak at 6 dpi, hereafter called profile 2. We next investigated the metabolic pathways sustained by the genes submitted to variation in stochasticity at these two stages. For this, we performed a Gene Ontology analysis of both groups of genes (Methods) and mapped the reactions they govern on the KEGG global metabolic map. As shown in Figure 6, the metabolic pathways corresponding to the genes allowing the functional annotation clustering of entropy kinetic profiles 1 and 2 cover a great deal of the murine metabolic map and hardly overlap (See also SupplementaryTable 5). Different sections of CD8 T cell metabolism are thus affected by the early and late increases in gene expression variability.

### Metabolism analysis reveals a transition stage 4 days post-infection

The late surge in gene expression inter-cellular variability suggests a new transition step at 4 dpi that may be responsible for the heterogeneity in fluxes and metabolite concentrations observed at that time of the response (Figure 2, Figure 4). It is indeed striking that the dichotomy observed at 4 dpi in module flux values and metabolite concentrations largely overlaps the clustering obtained based on all hypervariable gene expression (Methods, Figure 7A and data not shown). Furthermore, when cells collected at 4 dpi are clustered based on flux module values or metabolite concentrations, a large overlap is again observed between these clusters and those based on all hypervariable gene expression data (Figure 7B, C). Thus, the late surge in inter-cellular metabolic gene expression variability (Figure 5B, C) reveals a transition stage affecting all gene expression data and specifically metabolic activity, and points toward the existence of two distinct cell populations at that stage.

Gene set enrichment analysis (Methods) of D4-expression clusters mostly revealed an enrichment in genes associated with CD8 T cell effector functions (e.g., Ccl3, Ccl4, Ccl5, Ifng, Gzmb, Prf1) in cells from expression cluster #0 and the specific enrichment in “Zf5” motif of the Zbtb4 transcription repressor in 32 out of 47 over-expressed genes in cells from expression cluster #1 (SupplementaryTable 7).

## DISCUSSION

In this study we have made use of the scFEA algorithm (Alghamdi et al., 2021) to evaluate at the single-cell level the global metabolic activity in CD8 T cells responding to a viral infection *in vivo* from scRNA-seq data (Kurd et al., 2020; Milner et al., 2020). Previous single-CD8-T-cell metabolic analyses relied only on scRNA-seq measurements of mRNA levels of *in vitro* activated cells (Fernández-García et al., 2022), or cytometry quantification of a few selected



proteins implied in metabolism and/or its regulation (Ahl et al., 2020; Hartmann et al., 2021; Levine et al., 2021). However, mRNA and protein levels are not linearly correlated to the activity of the corresponding metabolic modules and our study is the first attempt at globally evaluating all metabolic activities in single responding CD8 T cells. The conclusions of our study can be questioned as they are drawn from flux value estimations from expression data. Two lines of evidence nevertheless support our claims: first, the output of the scFEA algorithm was validated against experimental data in the original study (Alghamdi et al., 2021) and second, our analysis does reproduce known facts on metabolic changes in activated CD8 T cells, demonstrating the validity of the approach. Indeed, flux estimation of carbohydrate metabolism pathways was in accordance with the extensively described switches in glucose consumption observed during CD8 T cell differentiation (van der Windt and Pearce, 2012; Klein Geltink et al., 2018; Gupta et al., 2020; Møller et al., 2022). Among the five modules describing the degradation of glucose to pyruvate, only M-2 (Glucose-6P (G6P) to G3P) showed a substantial flux variation but its flux increase upon activation is in agreement with a global rise in metabolic activity of antigen-challenged CD8 T cells. Furthermore, the flux increase in M-6 (pyruvate to lactate) exactly describes the metabolic switch from Oxphos to aerobic glycolysis that is crucial for CD8 T cell differentiation into effector cells. Oxphos is not entirely included in modules of the scFEA analysis, however the flux through M-8 (citrate to 2-oxoglutarate) and M-13 (malate to oxaloacetate) of the TCA, which fuels Oxphos, are decreased as expected. Of note, varying fluxes through modules may seem contradictory and sometimes difficult to interpret. However, metabolic stress analysis helped reconciling the observations on fluxomes. For instance, if the conflicting increase in M-11 (succinate to fumarate) is difficult to interpret directly in regards to M-8 and M-13 modules, metabolic stress analysis revealed a depletion of acetyl-coA that, together with the deprivation of G6P and pyruvate and the accumulation of lactate, all confirm the metabolic switch from Oxphos to aerobic glycolysis during the differentiation of virus-challenged CD8 T cells into effectors. Besides, the increased flux through M-62 (ornithine to putrescine) corresponds to the increase in polyamine synthesis from ornithine through the urea cycle, as described in Ag-activated T cells (Wang et al., 2011). scFEA analysis also added some temporal information on other known CD8 T cell metabolic perturbations. For instance, glucose consumption through the pentose phosphate anabolic pathway is increased in activated CD8 T cells (Sagone et al., 1974; Wang et al., 2011). It is thought to be important for nucleic acid demands in highly proliferating cells (Frauworth, Kenneth A. and Thompson, Craig B., 2004; van der Windt and Pearce, 2012). We observed that the flux through M-33 (G3P to PRPP) of the PPP initially decreases leading,

together with the increase flux through M-2, to an accumulation of G3P at 4 dpi. The increase in M-33 flux leading to an accumulation of PRPP at D6, is only seen secondary to the initial decrease, suggesting an important role for the PPP during the transition from activated cells to fully differentiated effectors. Similarly, T cell activation induces the increase of glutamine uptake and glutaminolysis (Carr et al., 2010; Wang et al., 2011). The glutamate produced by glutamine degradation can enter mitochondria and fuel the TCA to produce  $\alpha$ -KG and downstream cellular building blocks, such as lipids and nucleic acids (Gupta et al., 2020; Hansen and Gibson, 2022; Lane and Fan, 2015; Pearce et al., 2013), or be converted to glutathione in the cytoplasm to reduce the reactive oxygen species (ROS) produced by the enhanced mitochondrial activity (Klein Geltink et al., 2018; Møller et al., 2022). We observed an initial decrease in the flux through the M-25 (glutathione to glutamate), followed by an increase which led to accumulation of glutathione at D3 and glutamate at D5. This suggests that increased glutaminolysis is initially used to produce glutathione that will buffer the ROS production by mitochondria, in turn sustaining the mTOR pathway and the glycolytic switch (Mak et al., 2017). After 4 dpi, enhanced glutaminolysis leads to glutamate accumulation that can fuel the TCA cycle to produce *de novo* cellular components in highly proliferating effector cells (Gupta et al., 2020; Hansen and Gibson, 2022; Lane and Fan, 2015; Pearce et al., 2013). Finally, we detected the increased cholesterol production described in activated CD8 T cells (Chen et al., 1975; Kidani et al., 2013) only after 4 dpi, suggesting its role may not be major for the immune synapse formation (Yang et al., 2016) but important for the regulation of the transcriptional activity of Liver X receptor (Bensinger et al., 2008) and Sterol Regulatory Element-Binding Protein (Kidani et al., 2013), in activated T cells (Møller et al., 2022). In conclusion, scFEA analysis provided a very detailed description of metabolic adaptations in virus activated CD8 T cells at the single cell level that were validated by previous experimental results but added a level of temporal precision.

Nevertheless, no substantial variations of flux or metabolite concentration in cells were observed past 10 dpi and, thus, the back-switch in metabolism from aerobic glycolysis to Oxphos and FAO of effectors differentiating into memory CD8 T cells (Klein Geltink et al., 2018; Gupta et al., 2019; Møller et al., 2022) was not evidenced by our analysis. This may be due to variations in flux and metabolite concentrations that are beyond the sensitivity of scFEA estimations. Furthermore, fluctuation of fluxes in modules from ‘beta-alanine metabolism’ SM6, ‘urea cycle’ SM9, ‘spermine metabolism’ SM11 and ‘hyaluronic acid synthesis’ SM13 did not result in detectable accumulation or deprivation of any metabolite on the corresponding pathways, suggesting a reduced sensitivity of scFEA estimations of metabolite concentrations

as compared to modules fluxes. Finally, scFEA could not evidence flux variations in any of the modules of the SM4, 5, 7, 8, 14, 15, 18 or 19, defined in Alghamdi et al. (2021) as ‘glycan, glycosaminoglycan, glutamate, glycogen, sialic and fatty acids, beta-alanine and aspartate metabolism or synthesis’. We cannot conclude whether these sections of CD8 T cell metabolism are less affected by cellular differentiation or whether the algorithm fails to evidence them. However, as many genes encoding enzymes responsible for reactions of some of these pathways, such fatty acid elongation, are submitted to a surge in expression variability, we suspect scFEA is not sensitive enough to catch all CD8 T cell metabolic perturbations, further strengthening the quantitative importance of the changes that were detected. In that respect, scFEA revealed variations of full sections of cellular metabolism that were not expected from experimental studies. For instance, scFEA revealed perturbations of the fluxomes in ‘N- and O-linked glycan synthesis’ SM16 and SM17 with deprivation of two types glycan backbones. Interestingly, enhanced glycolysis and glutaminolysis fuel an increase in O-linked GlcNAcylation of nuclear proteins upon T cell activation (Kearse and Hart, 1991; Swamy et al., 2016) that triggers the Nuclear Factor kappa-light-chain enhancer of activated B cells (NF- $\kappa$ B) signaling pathway (Ramakrishnan et al., 2013). The differential regulation of specific modules and metabolites in SM16 and SM17 highlighted by scFEA could help designing experiments to decipher which post-translational modifications are crucial for CD8 T cell antiviral responses. Furthermore, reduced flux in several modules of leucine and isoleucine degradation to succinyl-coA and acetyl-coA in SM9 resulted in the accumulation of both these amino acids and participated with aerobic glycolysis to the deprivation in acetyl-coA. Thus, scFEA suggests a role for these amino acids in CD8 T cell responses that remains to be experimentally investigated. Deprivation of acetyl-coA, the principal giver of acyl groups for post-translational modifications (Walsh et al., 2018) shall also deserve experimental investigation. Similarly, scFEA analysis revealed an increase in the flux through several modules of the ‘purine synthesis’ SM20, leading to the accumulation of deoxyadenosine, as well as a very complex interaction of fluxomes in modules and variations of metabolite concentrations in the ‘pyrimidine synthesis’ SM21. The role of specific intermediates and metabolic modules of nucleic acid biosynthesis in CD8 T cell activation is totally unexplored. In conclusion, scFEA analysis points out new sections of cellular metabolism that deserve experimental investigation in the context of CD8 T cell antiviral responses and memory development.

Cell differentiation has long been seen as an instructive process with genetic programs orchestrated by a set of master regulator genes, the transcription factors, being solicited in every single cell submitted to environmental perturbation (Davis et al., 1987; Herskowitz, 1989; Whyte et al., 2013). The recent breakthrough of single cell omics technologies has largely challenged this view (Kato and Igarashi, 2019). Indeed, stochasticity of gene expression at the single cell level (Raj and van Oudenaarden, 2008) translates to stochasticity in the differentiation process (Chang et al., 2008; Kalmar et al., 2009). In this Darwinian view, cell differentiation proceeds in two steps: stochastic gene expression in response to a stimulus creates a certain degree of transcriptional uncertainty, where individual cells can initiate different genetic programs, leading to a high degree of inter-cell variability in gene expression. The subsequent selection of fit cells allows the return to a homogeneous population of differentiated cells, in which a new stable state of the GRN has been established (Moris et al., 2016; Richard et al., 2016; Stumpf et al., 2017; Papili Gao et al., 2020; Dussiau et al., 2022; Parmentier et al., 2022; Toh et al., 2022).

Although this differentiation mechanism has been demonstrated in all biological models examined to date, including *in vivo* situations (Dussiau et al., 2022; Toh et al., 2022), we show here for the first time that it also affects metabolic genes during an *in vivo* lymphocyte response to a viral challenge. Given the importance of a proper metabolic activity for the differentiation of CD8 T cells in effector and memory populations during an antiviral response, this process is thought to be tightly controlled. However, our results show that, as many other differentiation processes, CD8 T cell responses benefit from a stochastic exploratory search of the gene expression space during their differentiation at least at the level of the metabolic functions.

This implies the existence of selection steps of cells with the best fit state at some points of the process. Our observations suggest that such a selection step may occur around 4 dpi. Indeed, at this stage a large fraction of cells seems to recover flux values and metabolite concentrations similar to those in unstimulated cells, as if unable to sustain the metabolic changes occurring in other cells between D3 and D5 of the response. Furthermore, in many cases the flux direction in modules, as well as the corresponding metabolite accumulation/deprivation in cells, reverts between D3 and D5 highlighting a metabolic switch at D4. In addition, kinetic patterns of gene expression variability revealed two groups of gene controlling different sections of the cellular metabolism, that show a surge in entropy either directly after infection, or later starting at 4 dpi suggesting a transition stage at that time of the response. This is strengthened by the observation that heterogeneity of metabolic activities in cells collected at 4 dpi matches different states of the global GRN. Altogether, these results suggest a transition stage accompanied by a metabolic

switch in activated cells around 4 dpi, when they start to differentiate into cytotoxic effectors, that would result in the selection of the best metabolically fit cells.

In a whole, our study allowed a global survey of cellular metabolism in CD8 T cell responding to a viral infection *in vivo*, at the single-cell level, highlighting metabolic perturbations beyond known switches revealed by previous experimental approaches. This study also evidenced a transition step around 4 dpi, accompanied by a metabolic switch during the differentiation of antigen-activated cells into full-blown effectors, adding one more piece of evidence linking metabolic activity variations and differentiation processes.

## METHODS

### Data collection and processing.

Sequencing reads were generated by Kurd et al. (Kurd et al., 2020) and Milner et al. (Milner et al., 2020). For these studies, the authors transferred P14 TCR-transgenic CD8 T cells, which recognize a Lymphocytic ChorioMeningitis Virus (LCMV) epitope, to histocompatible hosts that were acutely immunized the day after with 10<sup>5</sup> plaque-forming units of LCMV Armstrong. Single responding P14 CD8 T cells from the spleens of immunized hosts were sorted at different dpi and loaded into Single Cell A chips for partition into Gel Bead In-Emulsions in a Chromium Controller (10x Genomics). Single-cell RNA libraries were prepared according to the 10x Genomics Chromium Single Cell 3' Reagent Kits v2 User Guide and sequenced (paired-end) on a HiSeq 4000 (Illumina).

We used data collected at 0, 3, 4, 5, 6, 7, 10, 14, 21, 32 and 90 dpi and downloaded reads from GEO 'release 2020-10-09' (GSE131847) with fastqdump from the SRA-toolkit suite (<https://www.ncbi.nlm.nih.gov/sra>), using the --split-files option.

Quality control of reads was performed with fastp (Chen, 2023): we discarded reads of too low quality, trimmed the others and removed adapters before selecting reads with a phred quality score (Ewing and Green, 1998) of at least 30.

Reads were then aligned to the mouse genome assembly GRCm38 ([https://www.ncbi.nlm.nih.gov/assembly/GCF\\_000001635.20/](https://www.ncbi.nlm.nih.gov/assembly/GCF_000001635.20/)) with the Kallisto-Bustools wrapper (Melsted et al., 2021), using the 'lamanno' workflow to generate a table of Unique Molecular Identifier (UMI) counts of spliced mRNAs. Empty sequencing droplets were removed with a threshold set to the inflection knee of the curve representing the UMI counts of

ranked droplets (<http://bioconductor.org/books/3.15/OSCA.advanced/droplet-processing.html>).

The UMI count tables of each collection day were imported and pooled in Seurat V4.0.4 (Hao et al., 2021) and the 14,666 genes that are expressed in at least 20 cells were selected. Dying cells with more than 5% of mitochondrial genes were further removed

The UMI count table was normalized with SCTransform (Hafemeister and Satija, 2019) and 2 % of the droplets containing more than one cell, as identified with the DoubletFinder package (McGinnis et al., 2019) were removed, resulting in an UMI count table of 14,666 genes expressed by 42,025 live single cells spanning the entire kinetics (Figure 1A). Principal Component Analysis (PCA) was performed with prcomp (<https://www.rdocumentation.org/packages/stats/versions/3.6.2/topics/prcomp>). The broken-stick model (MacArthur, 1957) was used to select 9 principal components for further dimension reduction and visualization with Uniform Manifold Approximation and Projection (UMAP).

### Metabolic gene selection

scFEA uses the expression of a list of 719 metabolic genes relevant to the estimation of the fluxes in all 171 metabolic modules. Among these metabolic genes, 430 are present in the 14,666 genes expressed in at least 20 cells in our data. However, all the genes regulating glycolysis and Oxphos described in Kyoto Encyclopedia of Genes and Genomes database (Kanehisa and Goto, 2000; Kanehisa et al., 2023, KEGG: release ‘101.4’) have not been included in the initial list. As metabolic switches from Oxphos to aerobic glycolysis and *vice versa* are crucial for the proper differentiation of CD8 T cells into effectors and memory cells during a viral infection, we complemented the list of 430 metabolic genes from N. Alghamdi et al. (2021) with 4 and 96 extra genes, respectively involved in glycolysis and Oxphos, and present in the 14,666 considered genes (SupplementaryTable 1). We thus filtered the raw UMI count table for the expression of the 530 selected metabolic genes. Interestingly, the expression level of many metabolic genes displays temporal variations during the CD8 T cell response (SupplementaryFigure 1).

### Data filtering and scFEA analysis

Raw UMI count data were filtered for metabolic genes in the designed list (SupplementaryTable 1) and the counts for 530 metabolic genes in the 42,025 cells were submitted to scFEA. The algorithm was installed and run at the High-Performance Computing cluster Pôle Scientifique de Modélisation Numérique of the Ecole Normale Supérieure de Lyon, according to the instructions of the authors (Alghamdi et al., 2021, <https://github.com/changwn/scFEA>).



## Module selection for fluxome analyses

The scFEA algorithm returned flux values in all cells for 168 modules. These were filtered to select fluxes with a  $|CV| > 7.10^{-4}$  for all cells to eliminate modules with too little flux variation along the kinetics (SupplementaryFigure 2). We checked the eliminated modules were not significantly varying during the CD8 T cell response and re-introduced the M-114 module that shows substantial variation with a  $|CV|$  of  $2.10^{-4}$  (SupplementaryFigure 2D).

To only consider relevant flux values, the 131 resulting varying modules were then filtered to select those with a maximum flux value superior or equal to  $10^{-3}$  AU. Among the 47 remaining modules, some still display weak flux values with a majority of cells not reaching the  $10^{-3}$  AU threshold (SupplementaryFigure 3A). They correspond to modules with a median of values inferior to  $3.10^{-4}$  AU (SupplementaryFigure 3B, C). These were filtered out and we finally removed from the 41 remaining modules, 7 modules corresponding to simple metabolite efflux from cells, resulting in a list of 34 modules of interest.

## Metabolite selection

The concentration of the 70 metabolites, for which scFEA yielded results, were averaged by day and the delta between the highest and the lowest concentrations along the kinetics (concentration\_delta) was calculated for each metabolite. The distribution of these concentration\_deltas displays a gap around  $10^{-3}$  AU (SupplementaryFigure 6) and we therefore selected the modules with a greater concentration\_delta, ending up with 20 metabolites of interest. However, concentration in ornithine was hardly varying along the response (Figure 4) and we did not consider it further, in the resulting top19 list (SupplementaryTable 3).

## Gene expression variability

The variability of gene expression is estimated by the level of entropy of each gene at each dpi (Gandrillon et al., 2021; Dussiau et al., 2022). Briefly, for each dpi, the distribution of gene expression among the cell population is binned and Shannon's entropy is then defined as minus the sum across bins of  $pk \cdot \log(pk)$ , where  $pk$  is the probability for a cell to belong to bin  $k$ . Shannon's Entropy thus measures heterogeneity in a population, with a value of 0 when all cells belong to the same bin (minimal entropy) and a maximal value of  $\log(k)/k$  when they are evenly distributed in bins (maximal heterogeneity). The entropy of each metabolic gene at each dpi was estimated with the unbiased 'best-upper-bound' estimator, as in Paninski et al. (Paninski, 2003).

## Clustering of entropy kinetic profiles

To identify groups of genes with similar temporal entropy patterns, i.e. similar profiles of entropies along the kinetics, we used functional PCA followed by kmeans (Ramsey, J. O. and Silverman, B. W., 2005). Briefly, functional PCA is used to reduce the dimensionality and catch temporal dynamics of entropy curves. We used a model with piece-wise constant functions (histograms) for their simplicity of interpretation. Then, a standard kmeans algorithm is used on principal components to find the appropriate clusters. The number of principal components and clusters ( $n = 2$ ) was determined by the rule of thumb.

## Gene Ontology analysis

The list of genes corresponding to the entropy kinetic profiles 1 and 2 in Figure 5C, were submitted to DAVID Bioinformatic Resources (<https://david.ncifcrf.gov/tools.jsp>) for functional annotation clustering on Gene Ontologies ‘Molecular Function’, ‘Biological Process’ and ‘Cellular Component’. Seventy-eight out of the 86 genes in entropy kinetic profile1 supported the clustering in 10 annotation clusters with low stringency and 107 out of the 444 genes in entropy kinetic profile2 supported the clustering in 15 annotation clusters with high stringency (SupplementaryTable 5). The 78 and 107 genes supporting both annotation clusters were then used to map the corresponding enzymatic reactions on the KEGG pathway map (Figure 6).

## Cell clustering

The 2,089 cells collected at 4 dpi were selected and the UMI counts of the 14,666 genes were normalized with SCTransform (Hafemeister and Satija, 2019). The selected 2,100 hypervariable genes were used for further dimension reduction with PCA and UMAP. Cell clustering was performed in Seurat (Hao et al., 2021) with a resolution level of 0.07 to obtain 2 clusters. Next, module flux values, as well as metabolite concentrations, in each D4-cell were normalized and clustered, with respective resolution values of 0.02 and 0.03 to again obtain 2 clusters for each clustering.

## Differential gene analysis and gene set enrichment

The 2,100 hyper-variable genes expressed by CD8 T cells at 4 dpi were selected and differential expression was evaluated by a kernel-based two-sample test that compares the distributions of gene expression (Ozier-Lafontaine et al., 2023). P-values were adjusted by the Bonferroni method. The 111 and 47 genes up-regulated in gene expression clusters #0 and #1, respectively, with an average  $\log_2(\text{FC}) > 0.13$  (SupplementaryTable 6) were then submitted to gene set

enrichment analysis in g:Profiler (<https://biit.cs.ut.ee/gprofiler/gost>) with default values and the 2,100 hypervariable genes used as the background list.

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## AUTHORS' CONTRIBUTIONS

Conceptualization, Methodology, Validation and Project Administration: C.A and O.G.; Software, Investigation, Data Curation and Writing: Original draft preparation: C.A., Formal Analysis: C.A. and F.P.; Writing: Review and editing: C.A., O.G. and F.P.; Supervision and Funding Acquisition: O.G.

## ABBREVIATIONS

ADP: Adenosine-DiPhosphate  
 Antigen: Ag  
 ATP: Adenosine-TriPhosphate  
 AU: Arbitrary Unit  
 coA: coenzyme A  
 CDP: Cytidine-DiPhosphate  
 CTP: Cytidine-TriPhosphate  
 dCDP: deoxyCytidine-DiPhosphate  
 dCMP: deoxyCytidine-MonoPhosphate  
 dCTP: deoxyCytidine-TriPhosphate  
 dpi: days post-infection

ETC: Electron transport Chain  
 FA: Fatty acid  
 FAO: FA oxidation  
 G3P: Glyceraldehyde-3-Phosphate  
 G6P: Glucose-6-Phosphate  
 GRN: Gene Regulatory Network  
 KEGG: Kyoto Encyclopedia of Genes and Genomes  
 LCMV: Lymphocytic ChorioMeningitis Virus  
 mTOR; mammalian Target Of Rapamycin  
 Oxphos: Oxidative phosphorylation  
 PCA: Principal Component Analysis  
 PPP: Pentose Phosphate Pathway  
 PRPP: 5-Phosphoribose-2P  
 ROS: Reactive Oxygen Species  
 scFEA: single-cell Flux Estimation Analysis  
 scRNA-seq: single-cell RNA-sequencing  
 SM: Super metabolic Module  
 TCA: TriCarboxylic Acid  
 TCR: T-Cell Ag Receptor  
 UMAP: Uniform Manifold Approximation and Projection  
 UMI: Unique Molecular Identifier  
 UMP: Uridine-MonoPhosphate  
 UTP: Uridine-TriPhosphate

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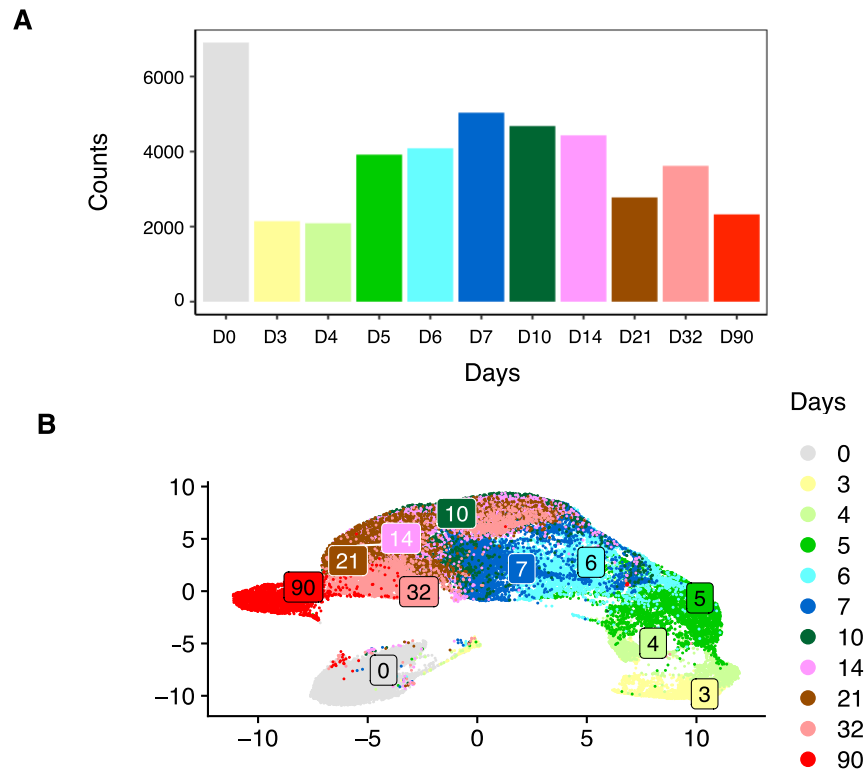
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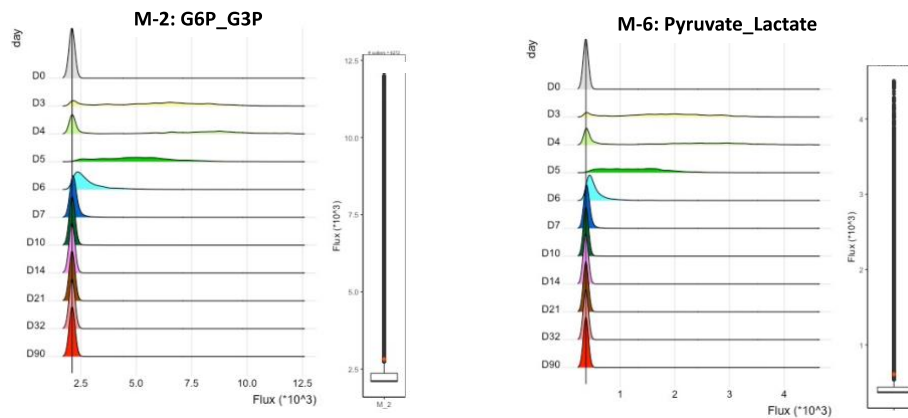
## FIGURES

**FIGURE 1: Data collection and processing.** (A) Bar plot of live single cell counts recovered after selection at the indicated dpi. (B) UMAP representation of the recovered cells.



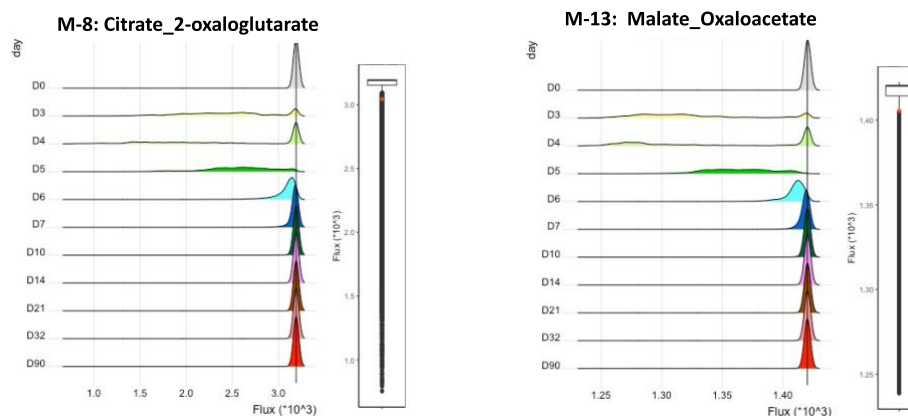
**FIGURE 2: Examples of module flux variations.** For each metabolic module, the distributions of flux values ( $\times 10^3$  AU) at all dpi are shown left-hand and a box-plot of flux values for all cells is shown right-hand. The vertical bar on the left and the red dot on the right indicate the median and mean of flux values, respectively. Examples of modules with an upregulated (A), downregulated (B) or biphasic (C) flux are shown. Names, as well as the initial and final compounds, of modules are indicated in the color of the corresponding Super-Module, as in Figure 3.

**A SM1: glycolysis**

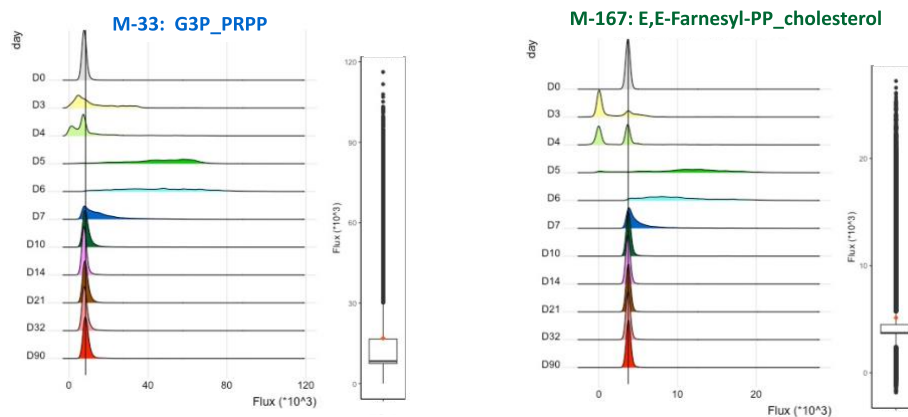


**B**

**SM1: TCA cycle**



**C SM3: pentose phosphate**





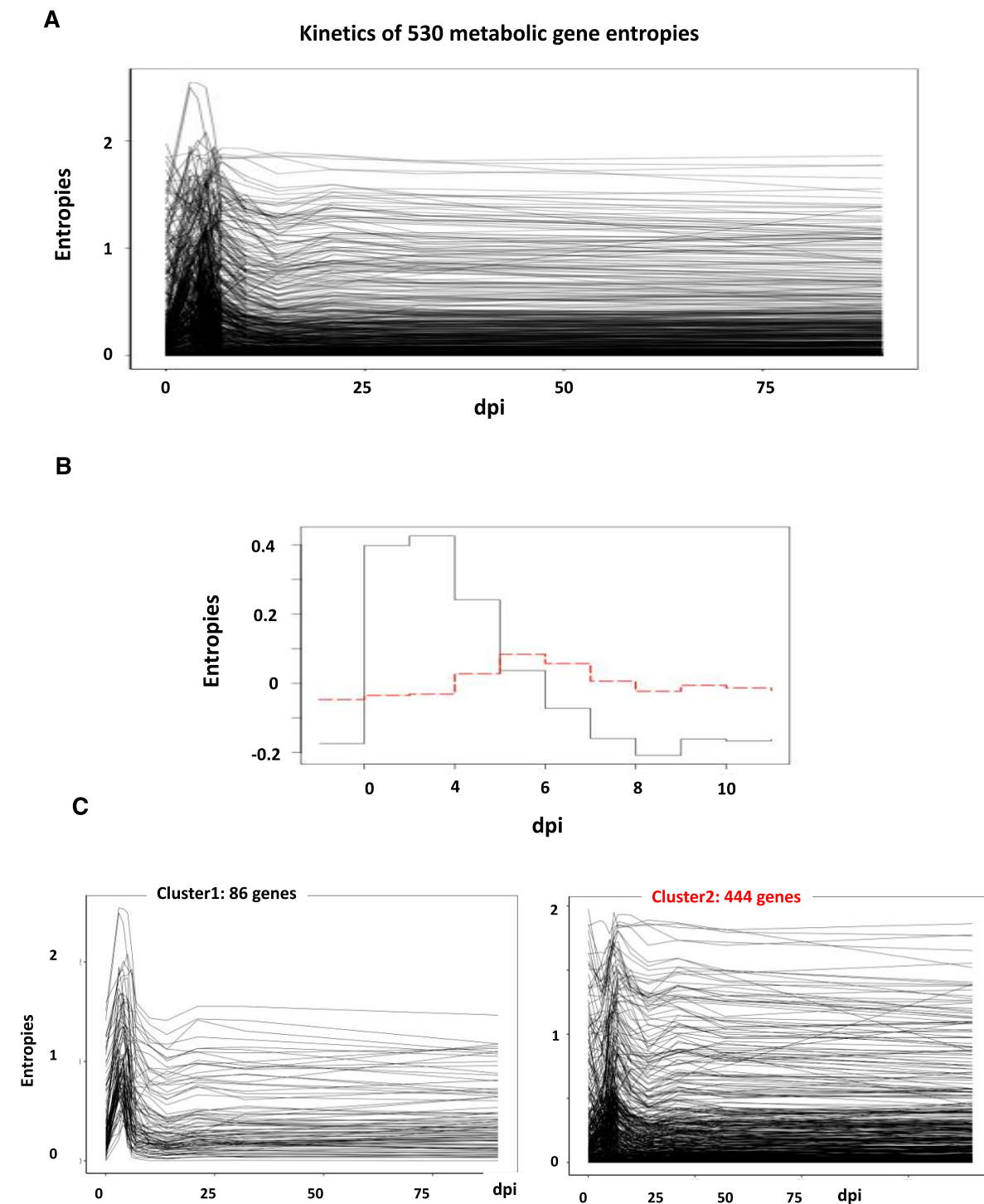




**FIGURE 4: Variation in concentration of the top20 most variables metabolites.** For each metabolite, the distributions of concentrations (AU) at all dpi are shown. The vertical bar indicates the median for all cells. Corresponding Super-Modules are indicated in their respective color. For metabolites at the junction of 2 SM, both are indicated.

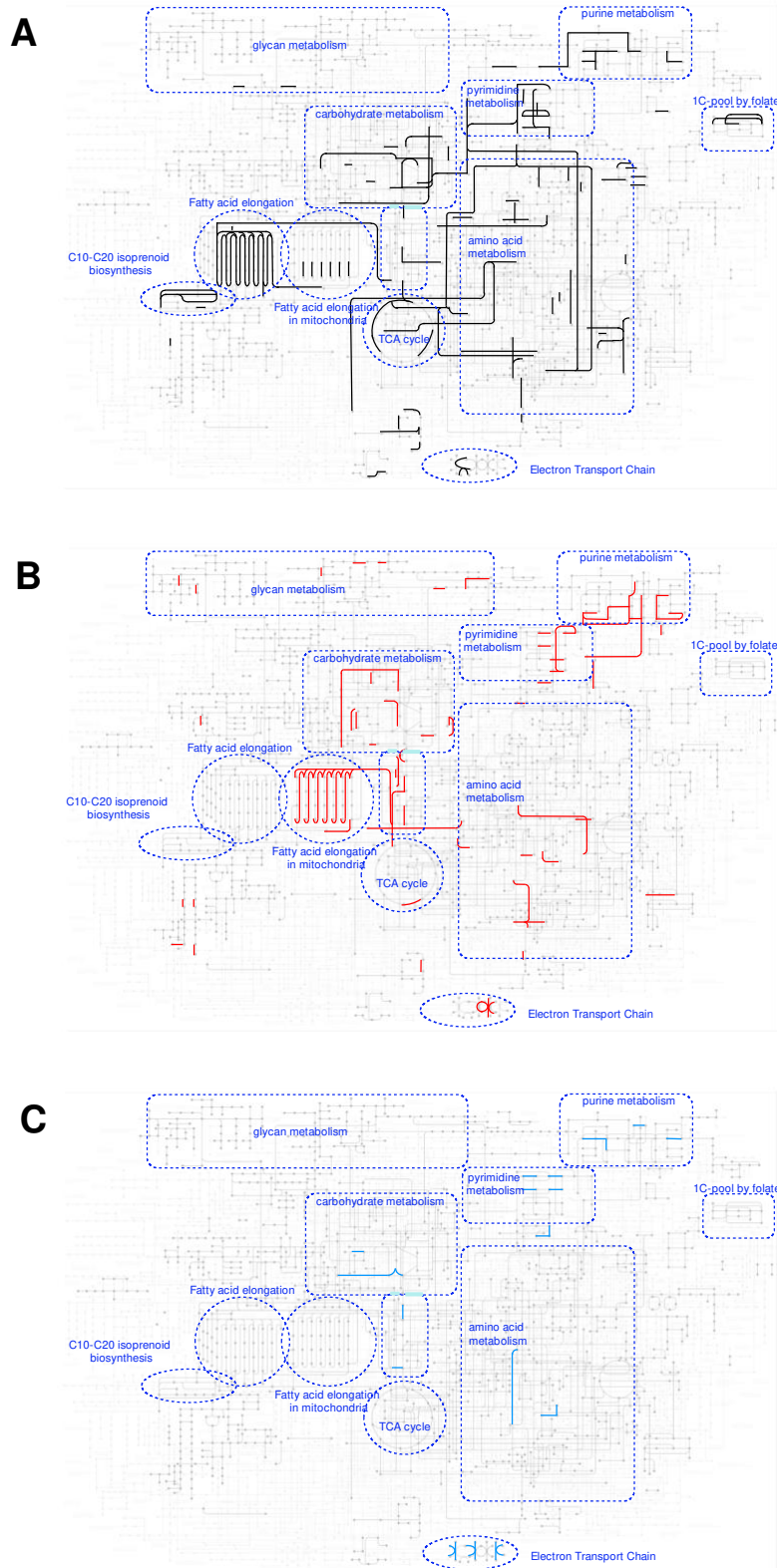


**FIGURE 5: Metabolic genes expression inter-cellular variability.** (A) The entropies of the 530 metabolic genes were calculated at each collection day and are represented as a function of time. (B) Functional PCA followed by kmeans clustering evidenced 2 groups of kinetic patterns. (C) The kinetics of the entropies of the genes in both clusters of (B) are shown as in (A).



# FIGURE 6: Metabolic map projection of the pathways covered by genes in kinetic profiles.

The enzymatic reactions corresponding to genes used for functional annotation of kinetic profiles are depicted on the KEGG murine metabolic map in (A) black for immediate profile 1, (B) red for later profile 2 and (C) blue for both. The regions corresponding to large pans of cellular metabolism are highlighted and labeled in blue.



# FIGURE 7: Module flux values and metabolite concentration of clustered D4 cells.

(A) Cells collected at 4 dpi were clustered on their gene expression levels and the histograms of flux values of two modules (top) and two metabolite concentrations (bottom) are shown according to the expression clusters #0 (red) and #1 (blue). (B) UMAP of D4 cell expression data colored by expression clusters (left, #0 in red and #1 in blue), flux module value clusters (middle, #0 in pink and #1 in green) and metabolite concentration clusters (right, #0 in orange and #1 in grey). (C) Cell repartition of flux module value clusters (left, #0 in pink and #1 in green) and metabolite concentration clusters (right, #0 in orange and #1 in grey) in expression data clusters.

