# An engineered biosensor enables dynamic aspartate measurements in living cells

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- Abstract Intracellular levels of the amino acid aspartate are responsive to changes in 10 metabolism in mammalian cells and can correspondingly alter cell function, highlighting the need 11 for robust tools to measure aspartate abundance. However, comprehensive understanding of 12 aspartate metabolism has been limited by the throughput, cost, and static nature of the mass 13 spectrometry based measurements that are typically employed to measure aspartate levels. To 14 address these issues, we have developed a GFP-based sensor of aspartate (iAspSnFR3), where the fluorescence intensity corresponds to appartate concentration. As a purified protein, the sensor has a 20-fold increase in fluorescence upon aspartate saturation, with dose dependent 17 fluorescence changes covering a physiologically relevant aspartate concentration range and no 18 significant off target binding. Expressed in mammalian cell lines, sensor intensity correlated with 19 aspartate levels measured by mass spectrometry and could resolve temporal changes in 20 intracellular aspartate from genetic, pharmacological, and nutritional manipulations. These data 21 demonstrate the utility of iAspSnFR3 and highlight the opportunities it provides for temporally 22
- <sup>23</sup> resolved and high throughput applications of variables that affect aspartate levels.
- 24

# 25 Introduction

The primary tool used by metabolism researchers, mass spectrometry (MS) coupled with either gas 26 chromatography (GCMS) or liquid chromatography (LCMS), involves extracting pools of thousands 27 of cells and measuring the liberated metabolites. This approach is powerful but has significant 28 drawbacks: it requires highly specialized equipment, it is expensive, and sample preparation by 20 chemical extractions homogenizes metabolic differences that may occur amongst different cells in 30 complex samples or across subcellular compartments. Metabolite extraction also consumes pre-31 cious samples that might otherwise be desirable to analyze over time or with additional outputs. 32 Development of genetically encoded protein sensors (biosensors) over the past two decades has 33 provided new opportunities to visualize the release, production, and depletion of important sig-34 naling molecules and metabolites with subsecond and subcellular resolution (reviewed in Kostvuk 35 et al. (2019); Koveal et al. (2020)). Thus, with the trade-off of only monitoring one metabolite per 36 sensor, biosensors provide a solution to many of the problems inherent to metabolite extraction 37 and MS. Aspartate is amongst the most concentrated metabolites in cells (Park et al., 2016), vet it is 39

Aspartate is amongst the most concentrated metabolites in cells (*Park et al., 2016*), yet it is one of only two amino acids that is not predominantly acquired from the environment. While the

- other, glutamate, is made from glutamine by the enzyme glutaminase, no analogous enzyme exists
- <sup>42</sup> in humans to convert asparagine to aspartate (*Sullivan et al., 2018*). Instead, aspartate must be
- 43 synthesized by transamination of the tricarboxylic acid (TCA) cycle metabolite oxaloacetate by the
- cytosolic enzyme GOT1 or the mitochondrial enzyme GOT2. Notably, aspartate synthesis can oc-
- 45 cur from multiple metabolic sources *via* complex metabolic reactions occurring in both the cytosol
- and mitochondria, rendering aspartate levels at the whole cell and subcellular levels dependent
- on multiple metabolic variables. For example, impairments to mitochondrial respiration can de-
- <sup>48</sup> plete aspartate levels and aspartate restoration can reestablish proliferation in cells with defective
- <sup>49</sup> mitochondria (Sullivan et al., 2015; Birsoy et al., 2015; Cardaci et al., 2015; Hart et al., 2023). Alter-
- ations to aspartate levels are associated with modifications to cell function in multiple biological
- processes, including stem cells (Tournaire et al., 2022; Arnold et al., 2022), immune cells (Bailis
- et al., 2019), endothelial cells (Diebold et al., 2019), and cancer (Helenius et al., 2021). In addition,
- <sup>53</sup> genetic methods to elevate intracellular aspartate can impact biology *in vivo*, increasing tumor <sup>54</sup> growth (*Sullivan et al., 2018*; *Garcia-Bermudez et al., 2018*) and improving hematopoietic function
- (*oi et al., 2021*). Therefore, our understanding of metabolism in multiple biological systems could
- <sup>56</sup> be improved with the availability of an aspartate biosensor.
- <sup>57</sup> We have previously developed a biosensor for glutamate (iGluSnFR) using the *E.coli* glutamate/aspartate
- <sup>58</sup> binding domain (GltI) linked to circularly permutated GFP (*Marvin et al., 2013*), and subsequently
- optimized it by modulating its affinity, kinetics, color, and total fluorescence change (SF-iGluSnFR
- and iGluSnFR3) (Marvin et al., 2018; Aggarwal et al., 2023). Since the GltI domain also binds aspar-
- tate, albeit at lower affinity than glutamate (*Hu et al., 2008*), we reasoned that subtle modifications
- $_{62}$  to the ligand binding site could switch the relative aspartate/glutamate specificity. We achieved this
- using a small mutagenesis screen on a precursor to iGluSnFR3 (Supplementary file 1), guided by
- the crystal structure of glutamate-bound GltI. The resulting biosensor, jAspSnFR3, was character-
- ized *in vitro* and in cells with matched LCMS determined aspartate levels, showing that it accurately
- reports genetic, pharmacological, and nutritional manipulation of intracellular aspartate.

## 67 Results

# **BROTE Protein engineering**

- $_{\tt 69}$   $\,$  We observed that the glutamate sensor, iGluSnFR, binds both glutamate and aspartate, with higher
- <sup>70</sup> affinity for the former (*Marvin et al., 2013*). To shift the relative affinities of the two ligands, we
- <sup>71</sup> evaluated the structure of the binding pocket (*Hu et al., 2008*), and sampled all possible amino acid
- <sup>72</sup> substitutions of residue S72, which interacts with the side-chain carboxylate of bound glutamate
   (*Figure 1*, panel A). By expressing mutant sensors in bacteria and measuring the fluorescence of
- <sup>73</sup> (*Figure 1*, panel A). By expressing mutant sensors in bacteria and measuring the fluorescence of bacterial lysate in response to aspartate and glutamate, we identified S72A and S72P as having
- $_{75}$  switched specificity from glutamate to aspartate. S72T, identified in a faster version of iGluSnFR
- 75 Switched specificity from glutamate to aspartate. 5721, identified in a faste
   76 (*Helassa et al., 2018*), also preferentially binds aspartate over glutamate.
- As an improved glutamate sensor (iGluSnFR3) was being developed (*Aggarwal et al., 2023*),
- we took a variant from that process and queried the effect of S72A, S72T, and S72P on aspar-
- tate/glutamate affinity. In bacterial cell lysate, S72P maintained the expected shift to a preference
- so for aspartate when inserted into a iGluSnFR3 precursor, and had a higher fluorescence fold in-
- <sup>81</sup> crease (F/F) than either S72A or S72T (*Figure 1—figure Supplement 1*, panel A). To further increase
- specificity of the S72P mutant, we sampled mutations at S27, which also interacts with the carboxy late of bound glutamate. One of those, S27A, had lower affinity for glutamate while mostly main-
- Iate of bound glutamate. One of those, S2/A, had lower affinity for glutamate while mostly maintaining affinity for aspartate (*Figure 1—figure Supplement 1*, panel A). We then moved forward
- taining affinity for aspartate (*Figure 1—figure Supplement 1*, panel A). We then moved forward with this variant, and since it is built from a precursor of iGluSnFR3, named it lanelia-developed
- Aspartate-Sensing Fluorescent Reporter (iAspSnFR3). Since we expected to be using this sensor
- in cell culture studies, and potentially *in vivo*, over the course of hours or even days, we added a
- <sup>88</sup> C-terminal red fluorescence protein. mRubv3, to enable correction for expression and movement
- artefacts. All biochemical characterization is reported with jAspSnFR3-mRuby3. For jAspSnFR3 sig-

<sup>90</sup> nal normalization in cells we used a mix of jAspSnFR3-mRuby3 and nuclear localized RFP.

Further characterization of the sensor found it is yellow-shifted in excitation and emission com pared to typical GFP-based sensors, since its chromophore is formed by the triad of GYG and has
 the T203Y pi-stacking mutations of the Venus yellow fluorescent protein. This yellow-shift facili-

- tates observation deeper into tissues using 2-photon microscopy, as the sensor has high signal fold
- change and significant 2-photon cross-section at 1040 nm (*Figure 1—figure Supplement 1*, panel B).
- It has a  $K_D$  for aspartate of about 50  $\mu$ M and binds glutamate and asparagine with  $K_D$  greater than
- 5 mM (*Figure 1*, panel B). It also does not appreciably change its green fluorescence in response
   to other amino acids (*Figure 1—figure Supplement 2*, panel A). Surprisingly, the mRuby3 compo-
- nent responds to some amino acids at high millimolar concentrations, indicating a non-specific
- effect, potentially interactions with the C-terminal histidine tag (*Figure 1—figure Supplement 2*, panel B). Notably, this increase in fluorescence is still an order of magnitude lower than the green
- <sup>102</sup> fluorescence response and it occurs at amino acid concentrations that are unlikely to be achieved
- in most cell types. The sensor also does not respond to other decoys considered relevant to aspartate metabolism nor to relevant pharmacological treatments (*Figure 1—figure Supplement 2*.
- panel C). A recently described and concurrently developed biosensor for aspartate is reported
- to be adversely affected by temperatures higher than 30C, causing lower maximum F/F (*Hellweg*
- *et al., 2023*). Our aspartate sensor appears unaffected by temperature up to 37°C, with the same maximum F/F at 37°C as compared to 30C (*Figure 1—figure Supplement 2*, panel D). Like all cpGFP-
- based sensors, it is sensitive to pH but changes in fluorescence due to aspartate far exceed what
- one might expect from changes in fluorescence due to physiologically attainable changes in intra-

cellular pH (Figure 1—figure Supplement 2, panel E). To determine whether it had the potential to

serve as an aspartate biosensor in mammalian cells, we expressed jAspSnFR3 in H1299 cells along

- with nuclear-RFP. Expression of jAspSnFR3 had no obvious toxic effects and H1299 jAspSnFR3 cells
- had visible fluorescence in the green channel (*Figure 1*, panel C). As it is the primary substrate for
- aspartate production, glutamine removal is expected to deplete aspartate levels. Indeed, we found
- that 24 hours of glutamine withdrawal abolished GFP signal while leaving RFP unchanged. These findings therefore supported the further testing of jAspSnFR3 as a method to quantify aspartate
- <sup>118</sup> levels over time in live mammalian cells.

# <sup>119</sup> jAspSnFR3 reveals the temporal dynamics of aspartate limitation

Having shown that iAspSnFR3-mRuby3 protein can measure the concentration of aspartate *in vitro*. 120 we wanted to test the usefulness of the sensor in cells. To that end, we generated stable cell lines 121 with constitutive expression of iAspSnER3-mRubv3 or iAspSnER3 and nuclear localized REP, and generated single cell clones from each to yield cell lines with uniform expression. In each case, 123 we then normalized GFP sensor signal to RFP signal to control for expression differences within 12/ and across cell lines. We also noted that normalization with nuclear RFP and RFP fusion were 125 highly correlated, enabling iAspSnFR3 sensor applications where nuclear RFP labeling is desirable 126 e.g. for counting cells at multiple timepoints using live cell imaging (Figure 2—figure Supplement 1. 127 panel C). An important motivation for using a biosensor for tracking aspartate changes is to enable 128 temporal measurements on the same subset of live cells, therefore we used an Incucyte S3 which 120 performs live cell imaging under native cell line growth conditions. 130

Cellular aspartate levels depend on the availability of metabolic precursors and the activity 131 of several metabolic processes. One such process is the generation of a sufficiently large intra-132 cellular NAD+/NADH ratio to drive aspartate precursor synthesis, a process normally maintained 133 through mitochondrial respiration or, in its absence, by treatment with exogenous electron ac-134 ceptors like pyruvate (Figure 2, panel A). Genetic alterations and pharmacological treatments that 135 disrupt mitochondrial respiration can decrease NAD+/NADH and aspartate levels, both of which 136 can be partially restored by supplementation with pyruvate (Sullivan et al., 2015: Birsoy et al., 137 2015). We thus tested the ability of iAspSnFR3 to quantify depletion of intracellular aspartate abun-138 dance upon treatment with the mitochondrial complex I inhibitor rotenone and the partial rescue 13



**Figure 1.** Protein engineering and *in vitro* characterization. (A) Structure of the binding pocket of glutamate-bound Gltl (2VHA.pdb) with residues S72 (left) and S27 (right) shown as sticks and bound glutamate as sticks inside transparent spheres. (B) Fluorescence response of purified jAspSnFR3-mRuby3 when titrated with aspartate (black) or glutamate or asparagine (grey tones). Ex. 485 nm (20 nm bandpass), Em. 535 nm (20 nm bandpass). Error bars are s.d. of three technical replicates. (C) Live cell imaging in the phase contrast, GFP and RFP channels of H1299 Nuclear-RFP cells expressing jAspSnFR3 after 24 hours with/without glutamine.

**Figure 1—figure supplement 1.** Aspartate specificity and excitation/emission spectra.

Figure 1—figure supplement 2. Decoy, temperature and pH sensitivity.

of aspartate by supplementing cells with pyruvate. Titrating rotenone in H1299 cells, we observed
 a dose dependent decrease in sensor fluorescence with increased rotenone, corresponding to
 the expected decrease in aspartate synthesis capacity, and a partial restoration of fluorescence
 in cells co-treated with pyruvate (*Figure 2*, panel B). This observation was extended to different
 cell lines with different rotenone sensitivities, corroborating the observation of decreased sensor
 fluorescence upon rotenone treatment and rescue by pyruvate supplementation (*Figure 2*—*figure 3 Supplement 1*, panel A, B and D).

We next evaluated the ability of the sensor to measure changes in aspartate without requir-147 ing treatment with a mitochondrial inhibitor. To this aim, we used CRISPR/Cas9 to generate an 148 H1299 cell line with a double knockout (DKO) of the genes glutamic oxalacetic transaminases 1 149 and 2 (GOT1/2 DKO), which renders cells unable to synthesize aspartate and therefore dependent 150 on aspartate uptake from the media (Garcia-Bermudez et al., 2022). Using these H1299 GOT1/2 151 DKO cells, we titrated media aspartate and observed that sensor fluorescence decreased upon as-152 partate withdrawal, approaching a steady-state after approximately 11 hours that corresponded 153 to the aspartate availability in the media (Figure 2, panel C). We note that 10 mM media aspartate. 154 a higher concentration than any other amino acid in media, is still unable to rescue sensor sig-155 nal significantly above aspartate depleted media, confirming previous observations that aspartate 156 has poor cell permeability and often requires 20 mM aspartate or more to robustly contribute to 157 intracellular aspartate pools (Sullivan et al., 2018). 158

<sup>159</sup> Metformin has slower inhibitor kinetics compared to rotenone

Metformin is a commonly used diabetes treatment that has been shown to act as a mitochondrial complex l inhibitor (*Owen et al., 2000; El-Mir et al., 2000; Andrzejewski et al., 2014; Wheaton* 

- 162 et al., 2014) and can decrease intracellular aspartate levels in a dose responsive way (Gui et al.,
- 2016). Whereas rotenone is a lipophilic molecule that can cross the cell membrane and act rapidly,
- <sup>164</sup> metformin is hydrophilic and poorly permeable to most cells, resulting in comparatively delayed
- kinetics for metformin to decrease mitochondrial respiration in intact cells. As rotenone and met-
- formin are often used interchangeably as complex I inhibitors, we wondered whether they have
- an equivalent temporal effect on aspartate or if the delayed effects of metformin on mitochondrial
- inhibition would similarly delay its effects on aspartate levels. To test this, we treated cells with two
- doses each of rotenone and metformin with roughly equivalent aspartate lowering effects and fol-
- lowed the sensor signal over time (*Figure 2*, panel E). We observed that the aspartate depleting
- effects of metformin acted slower than rotenone, with 30 nM rotenone reaching steady-state after
- <sup>172</sup> 20h and 2 mM metformin reaching a similar sensor response after almost 40h. These data there-
- fore provide orthogonal confirmation of the differential kinetics of these drugs on cell metabolism
- and highlight the temporal opportunities enabled by measuring aspartate levels by iAspSnFR3.
- Asparagine salvage diverts glutamine consumption

In most cancer cell lines, intracellular aspartate is derived primarily from glutamine oxidation, thus 176 making glutamine depletion an entry point for affecting aspartate metabolism. It has previously 177 been reported that asparagine, a product of aspartate metabolism, becomes essential upon glu-178 tamine starvation (Pavlova et al., 2018; Zhang et al., 2014). We hypothesize that asparagine be-179 comes essential in these conditions because glutamine starvation decreases synthesis of aspar-180 tate, slowing asparagine production, and because asparagine supplementation spares aspartate 181 consumption, allowing it to be redirected into other essential fates. However, it has been difficult to 182 measure metabolic changes during glutamine limitation because continuous glutamine consump-183 tion during the course of the experiment will result in progressive glutamine depletion and further 184 developing metabolic effects. One solution to this problem is to measure the temporal changes 185 in appartate levels over the course of glutamine starvation, a possibility enabled by fluorescence 186 based measurements of aspartate using jAspSnFR3. Indeed, we found that full glutamine deple-187 tion has a rapid and drastic effect on sensor signal and that this effect was delayed by adding back 188 glutamine (Figure 2, panel F). Aspartate signal did not robustly correlate with the concentration of 189 glutamine in the media in the short term, but instead we found that higher amounts of glutamine 190 in the media delayed the time until aspartate depletion, presumably corresponding to the time at 191 which glutamine is fully depleted and unable to support further appartate synthesis. Furthermore, 192 we found that adding 1 mM asparagine delayed the decrease in sensor signal, suggesting that 193 when asparagine can be salvaged from the media it diverts glutamine consumption that would 194 otherwise be purposed for asparagine synthesis via aspartate consumption. We note that, as this data is produced in real-time, the method can be used to dynamically find the optimal sampling 196 times to measure and compare intracellular levels of all the metabolites relevant to glutamine 197 starvation using mass spectrometry. 198

## jAspSnFR3 signal correlates with intracellular aspartate concentration

It is an important requirement for an aspartate sensor that it reflects the intracellular concentra-200 tion of aspartate over a biologically relevant range for several cell lines. Reference points for the 201 intracellular aspartate concentration can be generated using metabolite extraction and LCMS, but 202 it is important to note that this technique reports the total amount of aspartate summed across all 203 compartments, which can differ in their aspartate concentration (*Chen et al.*, 2016). The LCMS de-204 rived concentration also does not reflect protein crowding, aspartate binding to enzymes, or other 205 factors that would affect the free aspartate concentration. Nevertheless, LCMS is the standard ap-206 proach in studying metabolism and has previously been used to correlate aspartate levels with cell 207 proliferation (Gui et al., 2016; Hart et al., 2023). Thus, we titrated mitochondrial inhibitors of com-208 plex I (rotenone and metformin) and complex III (antimycin A), with or without pyruvate rescue, in 200 three different cell lines and waited 24 hours until aspartate had reached near steady-state levels 210



**Figure 2.** JAspSnFR3 resolves temporal aspartate changes in live cells. (A) Overview of aspartate metabolism and the effect of glutamine depletion, mitochondrial inhibition, GOT1/2 knockout and pyruvate/asparagine supplementation. GLN, glutamine. ETC, electron transport chain. LDH, lactate dehydrogenase. OAA, oxaloacetic acid. ASP, aspartate. ASN, asparagine. For (B), (C), (E) and (F), sensor signal over time shown as RFP normalized jAspSnFR3 signal following various perturbations of live cells. All experiments shown are normalized to a pre-treatment scan, then treated with the specified drug or amino acid and scanned 30 min following treatment. Grey dashed lines indicate the time of treatment. (B) H1299 cells treated with a rotenone titration and rescued by co-treatment with pyruvate. (C) H1299 GOT1/2 double knockout cells grown in media with 40 mM aspartate, washed thrice in media without aspartate and then changed into media with a titration of aspartate. (D) Western blot verification of H1299 GOT1/2 double knockout. (E) H1299 cells treated with a titration of glutamine with or without 1 mM asparagine. Markers indicate the average using available well replicates and are superimposed on a bootstrapped 95% confidence interval colored using the same color code as the markers. AU, arbitrary unit.

Figure 2—figure supplement 1. Rotenone titration in different cell lines. Figure 2—figure supplement 2. Narrow range glutamine limitation.

<sup>211</sup> before conducting a final measurement of sensor fluorescence, followed by immediate metabo-<sup>212</sup> lite extraction and quantitative LCMS measurements of aspartate levels using isotope dilution. We

- then compared sensor signals to LCMS derived aspartate concentrations and fitted a Hill curve
- to infer the intracellular aspartate concentrations at half-maximum sensor signal (*Figure 3*). For
- all three cell lines, we observe a monotonically increasing relationship between sensor signal and
- intracellular aspartate concentration, covering around two orders of magnitude. We also observe
- no relationship between sensor signal and intracellular glutamate levels (Figure 3—figure Supple-
- 218 ment 1). These observations validate the utility of our sensor in a biologically relevant range of

<sup>219</sup> aspartate concentrations without interference from glutamate.

Interestingly, the intracellular aspartate concentrations at half-maximum sensor fluorescence

is more than 17 fold higher than the aspartate  $K_D$  determined by *in vitro* characterization of the

- sensor. While the numbers inferred for intracellular aspartate are only point estimates, it is highly
- <sup>223</sup> unlikely that they are inaccurate to this degree. We speculate that the apparent cytosolic aspartate
- concentration is likely lower than the total aspartate concentration summed across all compart-
- ments. This suggests that binding of aspartate by jAspSnFR3 is in competition with other proteins
- and highlights that another advantage of using a biosensor is that measurements are made relative
- 227 to their native environment.



**Figure 3.** JAspSnFR3 signal predicts LCMS measured intracellular aspartate concentration. A Hill equation with top and bottom asymptotes, midpoint and slope as free variables is fitted to the datapoints and shown by the black line. The intracellular aspartate concentration at the inferred half maximum of RFP normalized jAspSnFR3 signal is reported in the red inserts. (A) Rotenone, metformin and antimycin A titrations in H1299 cells. (B) Rotenone titration in HEK293t cells. (C) Rotenone and metformin titrations in HT1080 cells. Markers indicate a single well from which both LCMS and jAspSnFR3 data was collected. Replicate wells have identical color and marker shape. AU, arbitrary unit.

Figure 3—figure supplement 1. jAspSnFR3 signal does not correlate with glutamate concentration.

# 228 Discussion

<sup>229</sup> A biosensor for aspartate is an important step towards improved understanding of aspartate metabolism.

<sup>230</sup> We have shown that our jAspSnFR3 sensor can resolve temporal changes in intracellular aspartate

to answer questions that would be impractical using LCMS. In most studies involving aspartate

<sup>232</sup> metabolism, metabolite extraction is performed 6-16 hours after treatment with the implicit as-

sumption that this is enough time to reach metabolic steady-state. Using our sensor, we have

shown that the time to reach steady-state can be much longer and depends on the treatment. Fu-

<sup>235</sup> ture studies seeking to understand the effects of treatments affecting aspartate levels can there-

<sup>236</sup> fore use real-time measurements using jAspSnFR3 to determine when cells have reached steady-<sup>237</sup> state and then perform metabolite extraction for LCMS.

Recently, a concurrently developed aspartate sensor based on SF-iGluSnFR was reported by another group (*Hellweg et al., 2023*). That aspartate sensor started with the S72A mutation, but then included 6 additional mutations identified by a combination of targeted screening and deep

mutational scanning. Our design differs from the one by Hellweg et al. (2023) by 25 mutations 241 (Supplementary file 1). Notably, we achieved aspartate selectivity by generating only two muta-242 tions in the binding pocket of its precursor, and so the majority of these amino acid differences 243 derive from us building off the next generation precursor for iGluSnER3, which has increased F/F 244 compared to iGluSnFR. Indeed, the sensor described here appears to have a larger signal change 245 both *in vitro* and in live cells, although head-to-head comparisons have not been performed. Im-246 portantly, while the sensor from *Hellweg et al.* (2023) is adversely affected by temperature at 37°C our sensor is not affected, allowing us to perform cell culture experiments at standard incubation 248 conditions. Another difference is that our iAspSnFR3 sensor has a higher affinity for all three rel-240 evant ligands (aspartate, asparagine, and glutamate): however, we found no discernible effect of 250 treatment with 1 mM asparagine on sensor signal in cell culture experiments and found no cor-251 relation between intracellular glutamate concentration and sensor fluorescence across treatment 252 conditions (Figure 3—figure Supplement 1). Collectively, we conclude that the iAspSnFR3 aspar-253 tate sensor reported here has biochemical features that makes it ideal for measuring intracellular 254 aspartate levels in live cells. 255 In summary, we report a novel fluorescence based biosensor that enables dynamic measure-256 ments of aspartate. This tool is free of significant interference from relevant metabolites in phys-257 iological intracellular systems and can resolve changes in aspartate from diverse treatment con-258 ditions in live cells over time. This approach to measuring aspartate will also have advantages

<sup>259</sup> ditions in live cells over time. This approach to measuring aspartate will also have advantages
 <sup>260</sup> compared to LCMS based metabolomics, including enabling high throughput experiments to iden <sup>261</sup> tify variables that affect aspartate levels, such as testing the effects of a drug library on cells in
 <sup>262</sup> multiwell plates or using FACS based selection during genetic screens. Another potential use for

this sensor would be to dissect compartmentalized metabolism, with mitochondria being a critical target. Altogether, adoption of iAspSnFR3 to measure aspartate levels will therefore provide novel

opportunities to understand this critical node of cell metabolism.

# 266 Methods and Materials

# 267 Sensor engineering and screening

The starting template for iAspSnFR3 was a variant along the path of making iGluSnFR3 (sequence 268 information in Supplementary file 1). Site saturation mutagenesis at positions S72 and S27 was 260 achieved by the uracil template method (Kunkel, 1985). Mutant libraries (maximum theoretical di-270 versity of 20 each) were transformed into T7 express cells. Individual colonies were picked into a 271 96-well plate containing auto-induction media (Studier, 2005) and shaken at 30C for 18-24 hours. 272 then harvested by centrifugation. Cell pellets were resuspended in PBS and repelleted by centrifu-273 gation 5 times over, then frozen as pellets overnight. The frozen 96-well plate was thawed by addi-274 tion of room temperature PBS, agitated by vortexing to resuspend and lyse cells, and then pelleted 275 again. 100 µL clarified lysate was added to each of two black 96-well plates and its fluorescence was 276 measured. Aspartate or glutamate was added (final concentration 100 uM) and fluorescence was 277 measured again. Wells that had a higher F/F for aspartate than glutamate were isolated, titrated 278 with aspartate and glutamate, and sequenced. After confirming that S72P was the most selective variant for aspartate from a library of S72X, a library of S27X was made in the background of S72P. 280 The selection process was repeated, and S72P+S27A was identified as the "best" aspartate sensor 281 and named iAspSnER3, mRuby3 was subsequently cloned at the C-terminus and this construct was 282 named jAspSnFR3-mRuby3. 283

# 284 Protein expression and purification

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<sup>285</sup> For large scale protein expression and purification, jAspSnFR3-mRuby3 was transformed into T7

- express cells and a single colony was grown in 300 mL auto-induction media (*Studier, 2005*) at
- <sup>287</sup> 30C for 18 hours. Cells were pelleted by centrifugation at 6000g, resuspended in PBS and 1 M
  - NaCl and frozen. The resuspended cell pellet was thawed, sonicated on ice (5 sec on, 5 sec off,

- <sup>289</sup> 10 min), and centrifuged at 6000g to remove cellular debris. The lysate was further clarified by
- <sup>290</sup> centrifugation at 350,000g for 1 hour, and then purified by IMAC on a HisTrap FF column, with a
- <sup>291</sup> 2 mL/min flow rate and elution from 0 to 200 mM imidazole over 120 mL. Fluorescent fractions
- <sup>292</sup> were pooled, concentrated by ultrafiltration, and dialyzed in PBS to remove endogenously bound
- <sup>293</sup> ligands. Protein concentration was determined by alkaline denaturation, and measurement at
- A447 (Ext. Coeff. 44,000 M-1 cm-1).

# 295 Sensor biochemical characterization

- <sup>296</sup> In vitro fluorescence measurements were performed on a Tecan Spark plate reading fluorimeter at
- <sup>297</sup> 28C, with the exception of the controlled temperature measurement, in which a BioTek Cytation 5
- was used. Concentrated jAspSnFR3-mRuby3 protein was diluted to 0.2  $\mu$ M in PBS for all measure-
- <sup>299</sup> ments. Decoy amino acids and pharmacologues were purchased from Sigma-Aldrich and solvated <sup>300</sup> as 100 mM stocks in PBS, with the exception of rotenone, which was resuspended in DMSO. Titra-
- as 100 mM stocks in PBS, with the exception of rotenone, which was resuspended in DMSO. Litrations were performed by making serial dilutions (1:2) of the stock compound into PBS, and adding
- $_{302}$  10 µL of that to 100 µL of 0.2 µM protein solution. Fluorescence was measured before addition of
- <sup>303</sup> compound, and F/F was calculated as (F(treatment)-F(initial))/F(initial).
- <sup>304</sup> Two-photon cross sections were collected for 1 µM solutions of protein in PBS with or without 10
- <sup>305</sup> mM aspartate, excited by pulses from a mode-locked Ti:Sapphire laser (Chameleon Ultra, Coherent,
- <sup>306</sup> 80 MHz, 140-fs pulse width, 1 mW power at the focus). Emission was detected by an avalanche
- <sup>307</sup> photodiode (PDM Series, Micro Photon Devices) with a 550 nm filter (88-nm bandpass).

# 308 Cell culture

- Cell lines were acquired from ATCC (HEK293T, H1299, HT1080) and tested to be free from mycoplasma (MycoProbe, R&D Systems). Cells were maintained in Dulbecco's Modified Eagle's Medium
- (DMEM) (Gibco, 50-003-PB) supplemented with 3.7 g/L sodium bicarbonate (Sigma-Aldrich, S6297),
- 10% fetal bovine serum (FBS) (Gibco, 26140079) and 1% penicillin-streptomycin solution (Sigma-
- Aldrich, P4333). Cells were incubated in a humidified incubator at 37°C with 5% CO2.

# 314 Generation of nuclear RFP cell lines

- Nuclear RFP cell lines were generated using 1e5 transducing units of EF1A-nuclear RFP lentivirus (Cellomics Technology, PLV-10205-50) by spinfection. Cells were seeded at 50% confluency in 6 well dishes, lentivirus was added to fresh media with 8 µg/µL polybrene, then added to cells and followed by centrifugation (900g, 90 mins, 30°C). Two days after infection, cells were sorted for high RFP expression using fluorescence-activated cell sorting (FACS). High RFP cells were then expanded and single-cell cloned by limiting dilution, plating 0.5 cells/well on a 96 well plate. Plates were then screened for RFP expression and localization using Incucyte S3 (Sartorius) and a suitable clone
- 322 chosen, expanded, and used for all subsequent experiments.

# 323 Lentiviral production and stable cell line generation

iAspSnFR3 and iAspSnFR3-mRubv3 were first cloned into entry vector pENTR1A (Fisher, A10462) 324 using NEBuilder HiFI DNA Assembly Cloning Kit (New England BioLabs, E2621). These donor con-325 structs were then used to transfer their insert into destination vectors: pLX304-CMV-Blast (Ad-326 dgene, 25890), pLenti-CMV-Hygro (w117-1) (Addgene, 17454 a gift from Eric Campeau & Paul Kauf-327 man), or pLX304-CAG-Blast using LR Clonase II (Fisher, 11791100), pLX304-CAG-Blast was gener-328 ated in house by swapping the CMV promoter region of pLX304-CMV-Blast with a CAG promoter 329 provided on synthetic DNA (Integrated DNA Technologies). Each plasmid sequence was verified 330 by whole plasmid sequencing (Plasmidsaurus). Lentivirus was generated by co-transfection of 331 HEK293T cells with destination vector plasmid DNA and the packaging plasmids pMDLg/pRRE (Ad-332 dgene, 12251), pRSV-Rev. (Addgene, 12253) and pMD2.G (Addgene, 12259) using FuGENE transfec-333 tion reagent (Fisher, PRE2693) in DMEM (Fisher, MT10017CV) without FBS or penicillin-streptomycin. 334 The supernatant containing lentiviral particles was filtered through a 0.45 µM membrane (Fisher, 335

- $_{336}$  9720514) and was supplemented with 8  $\mu$ g/ $\mu$ L polybrene (Sigma, TR-1003-G) prior to infection. For
- infection, cells were seeded at 50% confluency in 6 well dishes and centrifuged with lentivirus (900g,
- <sup>338</sup> 90 mins, 30°C). After 24 hours the media was replaced with fresh media and after 48 hours cells
- $_{339}$  were treated with either 1  $\mu$ g/mL blasticidin (Fisher, R21001) or 150  $\mu$ g/mL hygromycin (Sigma-
- Aldrich, H7772-1G) and maintained in selection media until all uninfected control cells died. After
- selection, cells were expanded and single-cell cloned by limiting dilution, plating 0.5 cells/well us-
- ing 2-3 96 well plates. These clones were incubated until 10-30% confluency and screened for
- <sup>343</sup> high GFP and RFP signal using Incucyte S3 (Sartorius). The highest expressing monoclonal cells <sup>344</sup> were selected and further expanded on 6 well plates and again screened for fluorescence using
- were selected and further expanded on 6 well plates and again screened for fluorescence using the Incucyte. From this a single clone was chosen, expanded and used for all subsequent experi-
- the Incucyte. From this a single clone was chosen, expanded and used for all subsequent experiments. Different cell lines received different vector-sensor combinations: HEK293T cells were in-
- fected with pLX304-CAG-iAspSnFR3-mRubv3 (blasticidin). HT1080 with pLenti-iAspSnFR3-mRubv3
- (hygromycin) and HT1080, H1299 and H1299 GOT1/2 DKO cells expressing nuclear RFP were in-
- fected with pLenti-jAspSnFR3 (hygromycin).

# 350 Generation of GOT1/2 double knockout (DKO) cells

- Protocol and guide RNA generation was identical to that described in Hart et al. (2023). Briefly,
- three chemically synthesized 2'-O-methyl 3'phosphorothioate-modified single guide RNA (sgRNA)
- sequences targeting GOT1 and GOT2 were purchased (Synthego; *Table 1*). A pool of all six sgRNAs
- for GOT1 and GOT2 were resuspended in nuclease-free water, combined with SF buffer (Lonza,
- V4XC-2032), and sNLS-spCas9 (Aldevron, 9212). 200,000 H1299 cells were resuspended in the resulting solution containing ribonucleoprotein complexes (RNPs) and electroporated using a 4D-
- <sup>356</sup> sulting solution containing ribonucleoprotein complexes (RNPs) and electroporated using a 4D-<sup>357</sup> Nucleofector (Amaxa, Lonza). Nucleofected cells were then expanded and single-cell cloned by
- <sup>357</sup> Nucleofector (Amaxa, Lonza). Nucleofected cells were then expanded and single-cell cloned by <sup>358</sup> limiting dilution by plating 0.5 cells/well in a 96 well plate. Gene knockout was confirmed using
- western blots.

Table 1. CRISPR guides.

Gene	sgRNA sequence (5'-3')
GOT1	CAGUCAUCCGUGCGAUAUGC
	GCACGGAUGACUGCCAUCCC
	CGAUCUUCUCCAUCUGGGAA
GOT2	UUUCUCAUUUCAGCUCCUGG
	CGGACGCUAGGCAGAACGUA
	UCCUUCCACUGUUCCGGACG

# 🚥 Intracellular jAspSnFR3 measurements

Experiments were conducted in DMEM without pyruvate (Corning 50-013-PB) supplemented with 3.7 g/L sodium bicarbonate 10% dialyzed fetal boyine serum (FBS) (Sigma-Aldrich, F0392) and 1% penicillin-streptomycin solution. To start an experiment, cells were trypsinized (Corning, 25051CI). resuspended in media, counted using a coulter counter (Beckman Coulter, Multisizer 4) and seeded onto 24-well dishes (Nunc, 142475) with an initial seeding density of 50,000, 70,000, 70,000 or 150,000 cells/well for H1299, H1299 GOT1/2 DKO, HT1080 and HEK293T, respectively. After 24h 366 (H1299, HT1080, HEK293T) or 48h (H1299 GOT1/2 DKO) incubation, treatment was added and 367 plates moved into an Incucyte S3 (Sartorius) live cell imaging platform inside a humidified incu-368 bator at 37°C with 5% CO2, Rotenone (Sigma-Aldrich, R8875), metformin (Sigma-Aldrich, D150959) 360 and antimycin A (Sigma-Aldrich, A8674) treatments were spiked-in as 20x solutions in water and 370 the 2 mM pyruvate (Sigma-Aldrich, P8574) was added as 500x stock in water. For treatments with 371 varying media aspartate (Sigma-Aldrich, A7219) or glutamine (Sigma-Aldrich, G5792), wells were 372 thrice washed and filled with media deplete of the given amino acid, then it was added as a spike-373 in at the specified concentration from a 20x solutions in water. For plates receiving asparagine 374

(Sigma-Aldrich, A7094), this was added to 1 mM from a 20x solution in water, with vehicle wells 375 receiving water. Live cell imaging was performed on the Incucyte S3 using the GFP and RFP chan-376 nels with default exposure times. Images were processed using the associated Incucyte software 377 to subtract background, define areas of cell confluence and GEP/REP signal and extract the sum 378 of the fluorescence signal in these areas. The data for the GFP signal, RFP signal, GFP/RFP ratio 379 and confluence for each well at each timepoint was exported and used for further data processing using Python code. The iAspSnFR3 signal (GFP channel) was normalized to an RFP signal, either 381 as a stably expressed nuclear localized REP (Nuc-REP) or mRubv3 C-term fusion to iAspSnFR3. For 382 temporal measurements the first scan was made 30 min after treatment with subsequent scans 383 indicated on relevant plots. For some experiments a pre-treatment scan was made shortly prior to 38/ treatment to normalize the data to this point. For comparisons of near steady-state measurements 385 of GFP/RFP versus mass spectrometry based metabolite measurements, a single scan was made 386 24h after treatment, the plate was then quickly moved to ice and metabolite extraction performed 387 (see below). Another plate was processed in parallel for cell volume determination using a coulter 388 counter and averaging across three replicate wells. The normalized iAspSnFR3 signal as a function 380 of intracellular aspartate concentration, f(c), was fitted by a baseline shifted Hill curve: 390

$$f(c) = t + \frac{b-t}{1+(c/m)^s}$$

With t, b being the top and bottom of the curve, respectively, describing the upper and lower 391 asymptotes of normalized jAspSnFR3 signal. The curve slope is described by s, also known as Hill co-392 efficient, and the midpoint (m) describes the intracellular appartate concentration at half maximum 393 jAspSnFR3 signal. The curve parameters were fitted to the data using the Broyden–Fletcher–Goldfarb–Shann 394 (BFGS) algorithm with an upper bound constraint on the top of the curve of 1.2 times the maximum 395 observed normalized iAspSnFR3 signal in any of the conditions on the same plot. Note that this 396 curve is not intended to represent a mechanistic model of the binding kinetics, rather the purpose 397 is to infer a reasonable estimate of the intracellular aspartate concentration at half maximum jAsp-398 SnFR3 signal. 399

# 400 Metabolite extraction

For polar metabolite extraction, a plate was move to ice and the media was thoroughly aspirated.
For H1299 and HT1080 cells, wells were washed once with cold saline (Fisher, 23293184). For
HEK293T cells, washing was omitted due to weak cell adherence. Then, 1 mL 80% HPLC grade
methanol in HPLC grade water was added, cells were scraped with the back of a P1000 pipet tip
and transferred to Eppendorf tubes. Tubes were centrifuged (17,000g, 15 mins, 4°C) and 800 µL of
the supernatant containing polar metabolites was transferred to a new centrifuge tube and placed
in a centrivap until dry.

# <sup>408</sup> Intracellular amino acid concentration measurements by isotope dilution

Dried samples were reconstituted with 40 uL 80% HPLC grade methanol containing 5 uM U-13C. 400 U-15N labelled canonical amino acid mix (Cambridge Isotope Laboratories, MSK-CAA-1) and trans-410 ferred to vials for measurement by LCMS. The peak area for each amino acid was divided by its la-411 belled standard to derive the response ratio. The response ratio was then mapped to a calibration 412 curve to infer the amino acid concentration and finally the intracellular concentration was calcu-413 lated by correcting for each step introducing a dilution, including the use of the total cell volume. 414 To make the calibration curves a non-labelled amino acid mixture was made from an analytical 415 amino acid standard without glutamine and asparagine (Sigma-Aldrich, A9906-1ML) and added 416 glutamine (Sigma-Aldrich, 76523-100MG) and asparagine (Sigma-Aldrich, 51363-100MG) to match 417 the concentration of the other amino acids. Using this mix, three replicates of a 12 point 2-fold 418 dilution series was made with a max concentration of 500 µM and a volume per dilution of 40 µL. 419 These were placed in a centrivap until dry and reconstituted with 40 µL 80% HPLC grade methanol 420

- 421 containing 5 μM U-13C, U-15N labelled canonical amino acid mix (Cambridge Isotope Laboratories,
- 422 MSK-CAA-1) and transferred to vials for measurement by LCMS. The peak area for each amino acid
- was divided by its labelled standard to derive the response ratio, then the best fitting calibration
- 424 curves for each amino acid were chosen among either linear, power or a second-degree polyno-
- mial. Each calibration curve was manually inspected for proper fit and measurements below or
- <sup>16</sup> above the concentration range of the dilution series were discarded.

# 427 Liquid Chromatography-Mass Spectrometry (LCMS)

- Metabolite quantitation was performed using a Q Exactive HF-X Hybrid Quadrupole-Orbitrap Mass
   Spectrometer equipped with an Ion Max API source and H-ESI II probe, coupled to a Vanquish Flex
- <sup>430</sup> Binary UHPLC system (Thermo Scientific). Mass calibrations were completed at a minimum of ev-
- ery 5 days in both the positive and negative polarity modes using LTQ Velos ESI Calibration Solution
- (Pierce). Polar Samples were chromatographically separated by injecting a sample volume of 1 L
   into a SeQuant ZIC-pHILIC Polymeric column (2.1 x 150 mm 5 mM, EMD Millipore). The flow rate
- 433 Into a SeQuant ZIC-pHILIC Polymeric column (2.1 x 150 mm 5 mM, EMD Millipore). The flow rate 434 was set to 150 mL/min, autosampler temperature set to 10 °C, and column temperature set to 30
- <sup>435</sup> °C. Mobile Phase A consisted of 20 mM ammonium carbonate and 0.1 % (v/v) ammonium hydrox-
- ide, and Mobile Phase B consisted of 100% acetonitrile. The sample was gradient eluted (%B) from
- the column as follows: 0-20 min.: linear gradient from 85% to 20% B; 20-24 min.: hold at 20% B; 24-
- 438 24.5 min.: linear gradient from 20% to 85% B; 24.5 min.-end: hold at 85% B until equilibrated with
- 439 ten column volumes. Mobile Phase was directed into the ion source with the following parameters:
- sheath gas = 45, auxiliary gas = 15, sweep gas = 2, spray voltage = 2.9 kV in the negative mode or 3.5
- kV in the positive mode, capillary temperature = 300 °C, RF level = 40 %, auxiliary gas heater temper-
- ature = 325°C. Mass detection was conducted with a resolution of 240,000 in full scan mode, with
- an AGC target of 3,000,000 and maximum injection time of 250 msec. Metabolites were detected
- over a mass range of 70-850 m/z. Quantitation of all metabolites was performed using Tracefinder
- 445 4.1 (Thermo Scientific) referencing an in-house metabolite standards library using 5 ppm mass 446 error.

# 447 Data analysis and plotting

- All data processing, curve fitting, plotting and statistics for experiments involving jAspSnFR3 ex-
- pressed in cell lines was made using Python code and data available on Github: www.github.com/
- 450 krdav/Aspartate-sensor

# 451 Plasmid availability

452 Submitted to Addgene under article ID 28238106.

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**Figure 1—figure supplement 1.** (A) Switching specificity of the iGluSnFR3 precursor from glutamate to aspartate using S72X library (left) and S72P, S27X library (right). Titrations with aspartate (solid lines) and glutamate (dashed lines) in bacterial lysate. (B) Excitation and emission spectra of jAspSnFR3-mRuby3. Left, 1-photon spectra. Excitation wavelength was varied from 400 nm to 520 nm (7.5 nm bandpass) while observing emission at 535 nm (10 nm bandpass). Emission wavelength was varied from 535 nm to 600 nm (10 nm bandpass) while exciting at 510 nm (7.5 nm bandpass). Fluorescence was measured both in the absence (dashed lines) and presence of 10 mM aspartate (solid lines). Right, 2-photon cross-sections, also ± 10 mM aspartate, with an overlay of calculated F/F (green). Vertical bar indicates 1040 nm.



Figure 1—figure supplement 2. (A) jAspSnFR3-mRuby3 does not appreciably change its green fluorescence in response to other amino acids (alanine, phenylalanine, glycine, histidine (red line), isoleucine, leucine, methionine, proline, glutamine, arginine, serine, threonine, valine, or tryptophan). Insert with aspartate in black and glutamate/asparagine in grey for comparison. Ex. 485 nm (20 nm bandpass), Em. 535 nm (20 nm bandpass), 0.2 µM purified protein in PBS. (B) jAspSnFR3mRuby3 shows increased red fluorescence at millimolar concentrations of all amino acids, with apparent responses to histidine at 100  $\mu$ M (red trace). Ex. 587 nm (20 nm bandpass), Em. 662 nm (20 nm bandpass). (C) jAspSnFR3-mRuby3 does not respond to other decoys: citrate, lactate, pyruvate, malate, alpha-ketoglutarate, cis-aconitate, succinate, fumarate, or oxaloacetate (orange squares); nor to relevant pharmacological treatments: rotenone (green squares) or metformin. The small increase in fluorescence from rotenone is likely due to the scattering of a visibly turbid solution; rotenone has very low solubility in water. Ex. 485 nm (20 nm bandpass), Em. 535 nm (20 nm bandpass). (D) jAspSnFR3-mRuby3 is not adversely affected by temperature. Fluorescence as a function of aspartate titration at 23°C (light grey), 30°C (medium grey), and 37°C (black). Error bars are standard deviation of three technical replicates. (E) pH sensitivity of jAspSnFR3-mRuby3 (green component). Ex 485 nm (5 nm bp), Em 515 nm (10 nm bp). Error bars are standard deviation of 5 technical replicates. Solid line is with 3 mM aspartate, dashed line is without aspartate.



**Figure 2—figure supplement 1.** jAspSnFR3 temporal response after rotenone treatment. RFP normalized jAspSnFR3 signal change over time following rotenone treatment of live cells. Related to **Figure 2**, panel B; however, these experiments are not normalized to a pre-treatment scan. Grey dashed lines indicate the time of treatment and the first scan occurs 30 min after this. (A) HT1080 cells using nuclear RFP to normalize the jAspSnFR3 signal. (B) HT1080 cells using an RFP fused to jAspSnFR3 (jAspSnFR3-mRuby3) for normalization. (C) Comparison between the steady-state signal of (A) and (B) with a linear regression shown as a red dashed line to show that nuclear RFP and RFP fusion normalizations are equivalent. (D) HEK293t cells using an RFP fused to jAspSnFR3 for normalization. For plots (A), (B) and (D) markers indicate the average using available well replicates and are superimposed on a bootstrapped 95% confidence interval colored using the same color code as the markers. For plot (C) markers indicate the average using available well replicates and errorbars are drawn as +/- the standard deviation of the replicates. AU, arbitrary unit.



**Figure 2—figure supplement 2.** Aspartate depletes when glutamine is limiting. RFP normalized jAspSnFR3 signal change over time following glutamine depletion in H1299 cells. Identical to **Figure 2**, panel F but with fewer glutamine concentrations and more well replicates. AU, arbitrary unit.





