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Felix Renaudin, Lucie Orliaguet, Florence Castelli, François Fenaille, Aurelie Prignon, et al.. Gout and pseudo-gout-related crystals promote GLUT1-mediated glycolysis that governs NLRP3 and interleukin-1 $\beta$  activation on macrophages. *Annals of the Rheumatic Diseases*, 2020, 79 (11), pp.1506-1514. 10.1136/annrheumdis-2020-217342 . hal-03203907

**HAL Id: hal-03203907**

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**Official URL:** <https://doi.org/10.1136/annrheumdis-2020-217342>

**To cite this version:**

Renaudin, Felix and Orliaguet, Lucie and Castelli, Florence [et al.]. *Gout and pseudo-gout-related crystals promote GLUT1-mediated glycolysis that governs NLRP3 and interleukin-1 $\beta$  activation on macrophages.* (2020) *Annals of the Rheumatic Diseases*, 79 (11). 1506-1514. ISSN 0003-4967

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# Gout and pseudo-gout-related crystals promote GLUT1-mediated glycolysis that governs NLRP3 and interleukin-1 $\beta$ activation on macrophages

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## ABSTRACT

**Objective** Macrophage activation by monosodium urate (MSU) and calcium pyrophosphate (CPP) crystals mediates an interleukin (IL)-1 $\beta$ -dependent inflammation during gout and pseudo-gout flare, respectively. Since metabolic reprogramming of macrophages goes along with inflammatory responses dependently on stimuli and tissue environment, we aimed to decipher the role of glycolysis and oxidative phosphorylation in the IL-1 $\beta$ -induced microcrystal response.

**Methods** Briefly, an in vitro study (metabolomics and real-time extracellular flux analysis) on MSU and CPP crystal-stimulated macrophages was performed to demonstrate the metabolic phenotype of macrophages. Then, the role of aerobic glycolysis in IL-1 $\beta$  production was evaluated, as well in vitro as in vivo using <sup>18</sup>F-fluorodeoxyglucose positron emission tomography imaging and glucose uptake assay, and molecular approach of glucose transporter 1 (GLUT1) inhibition.

**Results** We observed that MSU and CPP crystals led to a metabolic rewiring toward the aerobic glycolysis pathway explained by an increase in GLUT1 plasma membrane expression and glucose uptake on macrophages. Also, neutrophils isolated from human synovial fluid during gout flare expressed GLUT1 at their plasma membrane more frequently than neutrophils isolated from bloodstream. Both glucose deprivation and treatment with either 2-deoxyglucose or GLUT1 inhibitor suppressed crystal-induced NLRP3 activation and IL-1 $\beta$  production, and microcrystal inflammation in vivo.

**Conclusion** In conclusion, we demonstrated that GLUT1-mediated glucose uptake is instrumental during the inflammatory IL-1 $\beta$  response induced by MSU and CPP crystals. These findings open new therapeutic paths to modulate crystal-related inflammation.

## INTRODUCTION

Monosodium urate (MSU) and monoclinic calcium pyrophosphate dihydrate (m-CPPD) crystals are responsible for gout and m-CPPD deposition diseases, respectively. Both crystals activate the innate immune system and induce recurrent and painful flares, which are interleukin (IL)-1 $\beta$ -driven

## Key messages

### What is already known about this subject?

- ▶ A switch of cell metabolism from oxidative phosphorylation to aerobic glycolysis in order to support energy demand is a hallmark of inflammatory phenotype of macrophages and multiple immune-mediated inflammatory diseases such as cancer or autoimmune diseases.
- ▶ Gout and pseudo-gout flare depend on monosodium urate (MSU) and calcium pyrophosphate crystal-induced interleukin (IL)-1 $\beta$  production, respectively, by macrophages.

### What does this study add?

- ▶ MSU and monoclinic calcium pyrophosphate dihydrate (m-CPPD) crystal stimulation leads to a metabolic reprogramming of macrophages in favour of aerobic glycolysis.
- ▶ MSU and m-CPPD crystal-induced NLRP3 inflammasome activation and IL-1 $\beta$  production by macrophages, as well as microcrystal-mediated inflammation in vivo, rely on a de novo glucose uptake through glucose transporter 1.

### How might this impact on clinical practice or future developments?

- ▶ This study demonstrated the key role of inflammatory cell metabolism and glucose availability in the inflammatory process of microcrystal-related pathology. It thus highlights a potential new therapeutic path for acute and chronic patients' arthritis care.

inflammations caused by macrophage-mediated neutrophil infiltration and activation in joints.<sup>1,2</sup> IL-1 $\beta$  production is a two-step process which can both be activated by MSU and m-CPPD crystals. The first step involves nuclear factor- $\kappa$ B and mitogen-activated protein kinase pathways,<sup>3</sup> leading to pro-IL-1 $\beta$  synthesis, and the second one relies on the nucleotide-binding oligomerisation domain (NOD)-like receptor family, pyrin (NLRP) domain-containing 3 inflammasome complex and

(<http://dx.doi.org/10.1136/annrheumdis-2020-217342>).

## Correspondence to

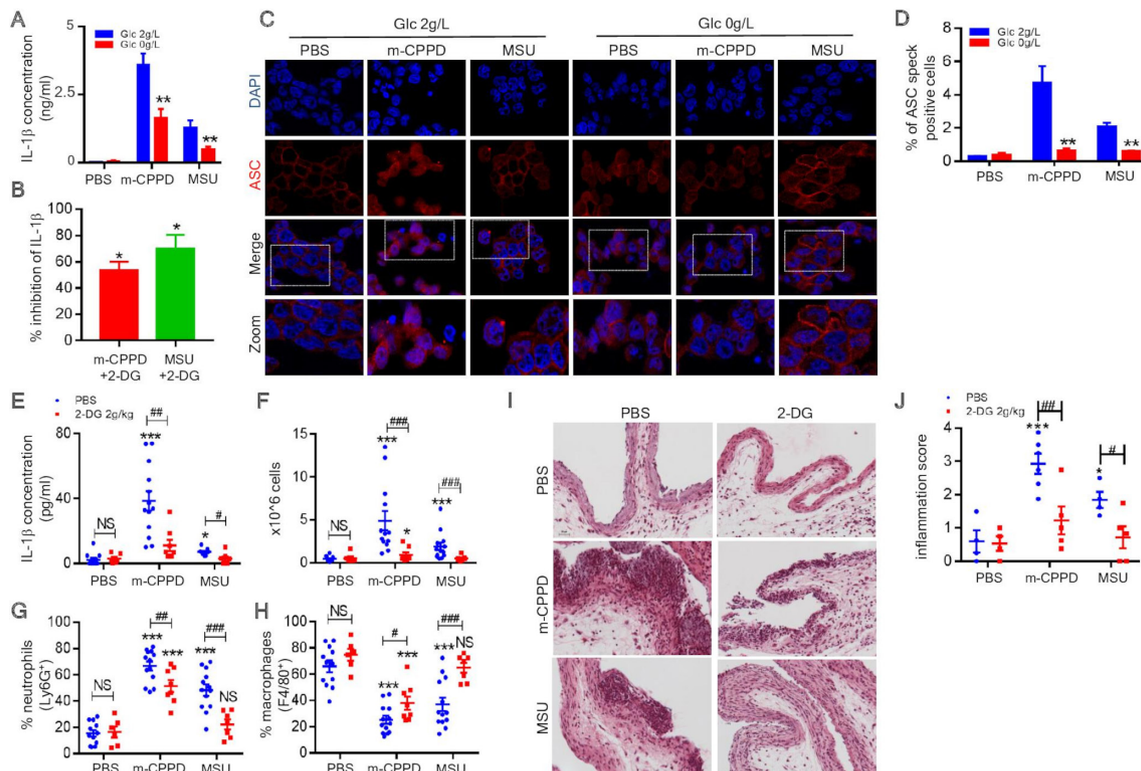
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To cite: Renaudin F, Orliaguet L, Castelli F, et al. *Ann Rheum Dis* Epub ahead of print: [please include Day Month Year]. doi:10.1136/annrheumdis-2020-217342

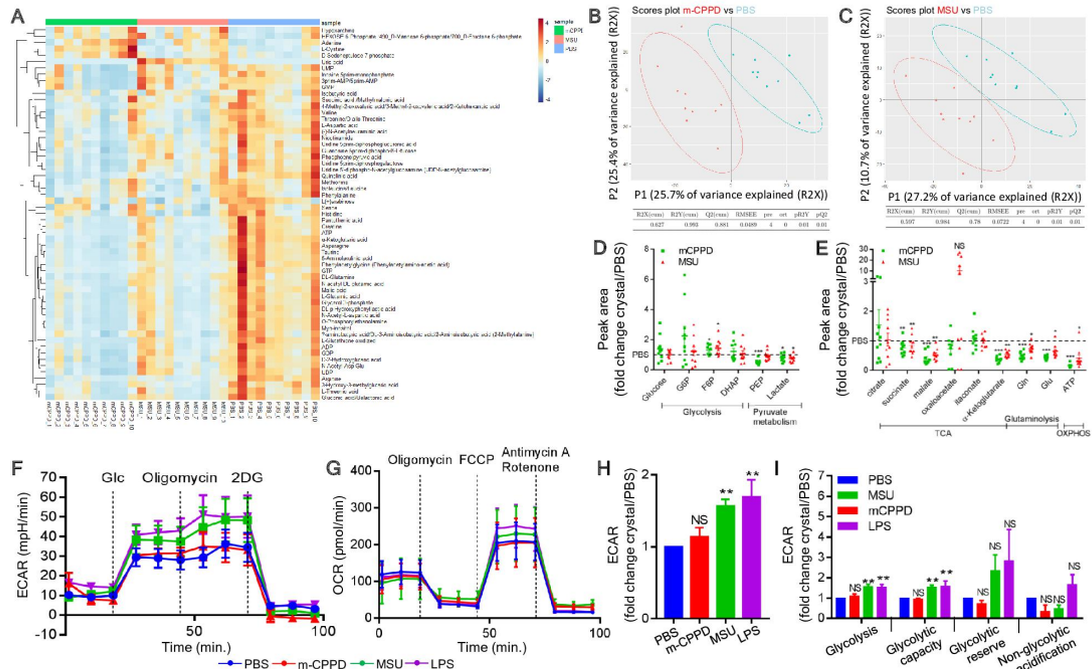
gives rise to the secretion of active IL-1 $\beta$ .<sup>1</sup> Activated NLRP3 stimulates speck formation of the adaptor protein apoptosis-associated speck-like containing a CARD (ASC) and recruits pro-caspase-1 into the NLRP3/ASC complex through homotypic domain–domain interactions.<sup>4,5</sup> Assembly of pro-caspase-1 within the NLRP3 inflammasome allows its autoproteolysis and the release of active caspase-1, which cleaves its substrates pro-IL-1 $\beta$  and pro-IL-18 into their mature forms.<sup>4</sup> NLRP3-deficient macrophages are unable to produce mature IL-1 $\beta$  under MSU and m-CPPD crystal stimulation.<sup>1,3</sup> Blocking IL-1 $\beta$  abrogates MSU-induced and m-CPPD-induced inflammation and constitutes an efficient therapeutic option in gout flare.<sup>6,7</sup> However, long-term inhibition of IL-1 $\beta$  may increase infection risk. Thus, understanding mechanisms of NLRP3 activation by MSU and m-CPPD crystals might offer a safer way to modulate IL-1 $\beta$  effects.

NLRP3 can be activated through several mechanisms, including reprogramming of cell metabolism.<sup>8–11</sup> First described in cancer cells, the so-called Warburg effect characterised by an increase in glucose uptake and aerobic glycolysis, along with a reduction of the mitochondrial respiration (oxidative phosphorylation (OXPHOS)), and an inhibition of tricarboxylic acid (TCA) cycle plays a critical role in host defence and inflammation and is a metabolic hallmark of activated immune

cells and proinflammatory macrophages.<sup>8–10</sup> Thus, high concentrations of glucose increase IL-1 $\beta$  production through the NLRP3-dependent pathway, while inhibition of glycolysis with 2-deoxyglucose (2-DG) suppresses IL-1 $\beta$  production by macrophages stimulated with the TLR-4 agonist lipopolysaccharide (LPS) or ATP.<sup>12,13</sup> LPS-induced glycolysis stimulates IL-1 $\beta$  production through hexokinase (HK) 1, hypoxia-inducible factor 1 $\alpha$  (HIF-1 $\alpha$ ) and pyruvate kinase muscle (PKM) 2 activation, three molecules of the glycolysis pathway directly involved in NLRP3 activation and IL-1 $\beta$  production.<sup>14–16</sup> In parallel, TLR4 activation induces TCA cycle alteration stimulating IL-1 $\beta$  production through cytosolic accumulation of succinate, which prevents degradation of HIF-1 $\alpha$  by prolylhydroxylase enzyme.<sup>12</sup> Stabilised HIF-1 $\alpha$  then enhances the expression of genes encoding IL-1 $\beta$  and proteins involved in glycolysis pathways, such as glucose transporter 1 (GLUT1) and HK, which further amplify glucose uptake and glycolysis.<sup>12,15,17</sup> Interestingly, glucose uptake quantified with <sup>18</sup>F-fluorodeoxyglucose ([<sup>18</sup>F]-FDG) positron emission tomography (PET) is increased in joints with gout flare and in soft-tissue surrounding MSU or m-CPPD crystal deposition, suggesting that glucose consumption plays an important role in crystal-induced inflammation.<sup>18,19</sup> Moreover, the ketone body  $\beta$ -hydroxybutyrate produced during starvation or low-carbohydrate ketogenic diet inhibits IL-1 $\beta$  production



**Figure 1** Crystal-induced inflammation depends on glucose availability. Primed THP-1 cells were stimulated with either PBS or MSU or m-CPPD crystals. (A,B) IL-1 $\beta$  production was quantified by ELISA in supernatants of cell culture in the (A) absence or presence of glucose (n=9). Multiple t-test with false discovery rate (FDR) correction between +glucose and –glucose (\*): #p<0.05, ##p<0.01, ###p<0.001. (B) In the absence (PBS) or presence of 2-DG (20mM, n=4). Data are presented in % of inhibition of IL-1 $\beta$  production compared with PBS (100%). Kruskal-Wallis test with FDR correction (\*): \*p<0.05, \*\*p<0.01, \*\*\*p<0.001. (C,D) ASC speck formation was observed with confocal microscopy and quantified in cells cultured with medium containing or not containing glucose. (C) Imaging representative of three independent experiments and (D) quantification of cells expressing at least one speck complex (n=3). Multiple t-test with FDR correction between +glucose and –glucose (#): #p<0.05, ##p<0.01, ###p<0.001. (E–J) Mouse air pouch model of microcrystal inflammation: IL-1 $\beta$  concentration (E), cell infiltration (F), proportion of neutrophils and macrophages (F,G) in the air pouch lavages of mice injected by either PBS, MSU or m-CPPD crystals and treated with 2-DG or PBS (n=12/group); H&E staining of air pouch membranes (I) and scoring of the inflammation (J) (n=5). Two-way analysis of variance test with FDR correction (#): #p<0.05, ##p<0.01, ###p<0.001. 2-DG, 2-deoxyglucose; ASC, apoptosis-associated speck-like containing a CARD; IL, interleukin; m-CPPD, monoclinic calcium pyrophosphate dihydrate; MSU, monosodium urate; NS, not significant.



**Figure 2** MSU and m-CPPD crystals induce metabolic changes in the glycolysis pathway and the TCA cycle. (A–E) Metabolomics analysis was performed by mass spectrometry in primed THP-1 cells treated by either PBS, MSU or m-CPPD crystals. (A) Heatmap analysis and (B,C) principal component analysis of MSU and m-CPPD crystals versus PBS (n=10). Relative quantification (ratio of MSU or m-CPPD to PBS) of the different metabolites related to glycolysis (D) and Krebs cycle or OXPHOS (E). Kruskal-Wallis test with FDR correction (\*): \* $p < 0.05$ , \*\* $p < 0.01$ , \*\*\* $p < 0.001$ . (F–H) Real-time extracellular flux analysis on LPS-primed BMDMs stimulated with either MSU or m-CPPD. (F) Time course of real-time changes in the ECAR after Glycolysis Stress Assay (n=6). (G) Time course of real-time changes in the OCR after Cell Mito Stress Assay (n=4). (H) Mean of basal ECAR (ratio of crystals over PBS). (I) Glycolysis Stress Assay (n=6): each rate was determined according to the SeaHorse Agilent Guide and described in the Materials and methods section. Kruskal-Wallis test with FDR correction (\*): \* $p < 0.05$ , \*\* $p < 0.01$ , \*\*\* $p < 0.001$ . 2-DG, 2-deoxyglucose; BMDM, bone marrow-derived macrophage; DHAP, dihydroxyacetone phosphate; ECAR, extracellular acidification rate; FCPP, carbonyl cyanide-4-(trifluoromethoxy) phenylhydrazone; LPS, lipopolysaccharide; m-CPPD, monoclinic calcium pyrophosphate dihydrate; MSU, monosodium urate; NS, not significant; OCR, oxygen consumption rate; OXPHOS, oxidative phosphorylation; PEP, phosphoenolpyruvate; TCA, tricarboxylic acid.

by macrophages and inflammation in a mouse gout model.<sup>20 21</sup> Nevertheless, whether macrophage metabolism reprogramming is involved in MSU and m-CPPD crystal-induced IL-1 $\beta$  production remains unknown.

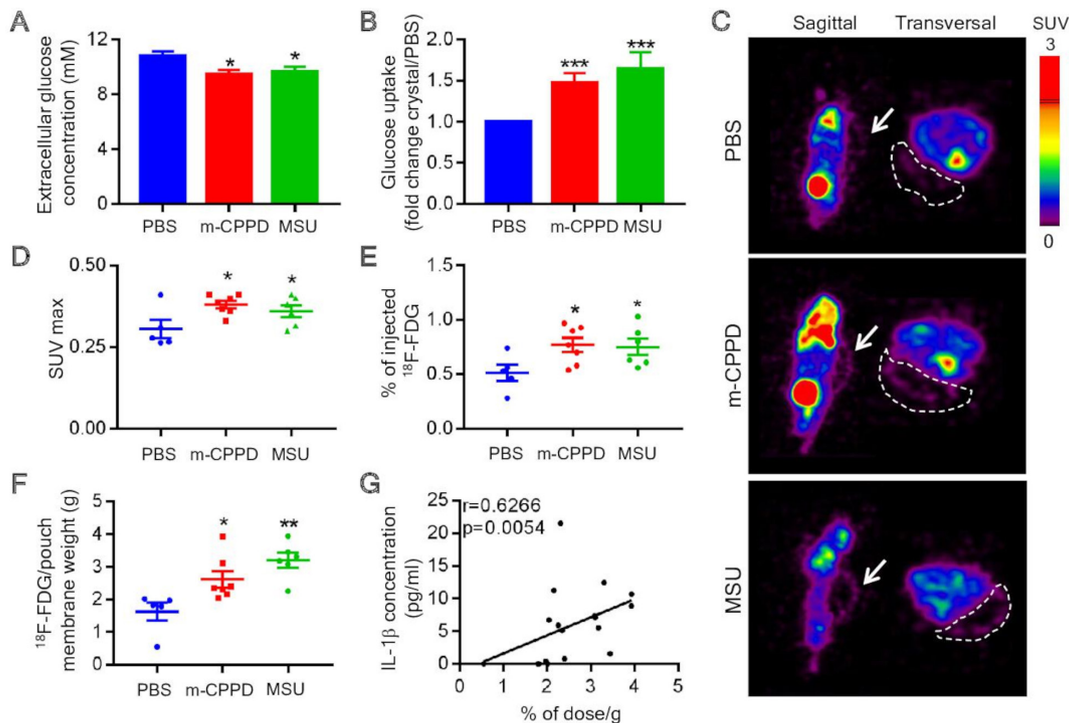
In this study, we aimed to assess the metabolic phenotype of macrophages and the role of glucose uptake in the NLRP3-dependent IL-1 $\beta$  production in response to MSU and m-CPPD crystals. We observed that MSU and m-CPPD crystals induced metabolic modifications in macrophages towards an increase in glycolytic activity. Upregulation of glycolysis corroborated with a de novo glucose uptake mediated by the glucose transporter (GLUT) GLUT1, in response to microcrystals. Interestingly, glucose deprivation or glycolysis inhibition by knock-down of GLUT1 prevented ASC oligomerisation (NLRP3 activation) and IL-1 $\beta$  secretion induced by both crystals. Moreover, both inhibition of GLUT1 and glycolysis inhibition by 2-DG decreased MSU and m-CPPD crystal-induced inflammation in an in vivo mouse model. Finally, in patients with gout flare, neutrophils isolated from the inflamed joint expressed more frequently GLUT1 at their surface membrane than circulating neutrophils, which highlights a promising specific approach for GLUT1 targeting as a gout flare therapy.

## RESULTS

### MSU and m-CPPD crystal-induced inflammation depends on glucose metabolism

To assess the role of glucose metabolism in crystal-induced inflammation, we stimulated cells with media containing

increasing concentrations of glucose (0–4 g/L) or supplemented with the glucose analogue 2-DG, an inhibitor of glycolysis. Both glucose deprivation and glycolysis inhibition drastically decreased IL-1 $\beta$  production by primed THP-1 cells or mouse bone marrow-derived macrophages (BMDMs) stimulated by either MSU or m-CPPD crystals (glucose deprivation decreased by more than 90% the crystal-induced IL-1 $\beta$  production by BMDMs) (figure 1A,B; online supplementary figure S1A,B). In contrary, increasing concentrations of glucose-enhanced IL-1 $\beta$  production (see online supplementary figure 1C). Nevertheless, prostaglandin E2 and CXCL1 production, two proinflammatory mediators involved in crystal inflammation, did not depend on glucose availability (see online supplementary figure S1E–F). Suppression of IL-1 $\beta$  production is secondary to the inhibition of NLRP3 inflammasome.<sup>13</sup> Here, we demonstrated that glucose deprivation suppressed ASC oligomerisation and speck formation induced by either MSU or m-CPPD crystals (figure 1C,D). We confirmed, in vivo, the central role of glycolysis and observed that mice treated with 2-DG displayed mild inflammatory response compared to untreated mice 6 hours after MSU or m-CPPD crystal stimulation. Indeed, 2-DG abrogated crystal-induced IL-1 $\beta$  production and prevented crystal-induced neutrophil infiltration assessed in the air pouch lavages (figure 1E–H). Histology analysis of air pouch membranes after H&E staining further evidenced that glycolysis inhibition suppressed neutrophil infiltration with a major decrease in inflammation score in mice treated by 2-DG (figure 1I,J).



**Figure 3** Crystal-induced inflammation is associated with an increase in glucose uptake. (A) Extracellular glucose concentration in cell culture media was quantified 6 hours after crystal stimulation ( $n=5$ ). (B) [ $^{18}\text{F}$ ]-fluorodeoxyglucose ([ $^{18}\text{F}$ ]-FDG) was quantified in cell pellets to assess glucose uptake after 1 hour of stimulation by either MSU or m-CPPD crystals ( $n=8$ ). (C–E) Glucose uptake was quantified in air pouch (arrow) model using [ $^{18}\text{F}$ ]-FDG PET/CT. (C) Imaging representative of 6 mice; (D)  $\text{SUV}_{\text{max}}$  of air pouch; and (E) proportion of injected [ $^{18}\text{F}$ ]-FDG dose detected in the air pouch ( $n=6$  mice per group). (F) Ratio of [ $^{18}\text{F}$ ]-FDG per weight (g) of ex vivo isolated air pouch membranes. Kruskal-Wallis test with FDR correction (\*): \* $p<0.05$ , \*\* $p<0.01$ , \*\*\* $p<0.001$ . (G) Correlation between IL-1 $\beta$  concentration (pg/mL) in the air pouch and the [ $^{18}\text{F}$ ]-FDG quantification (% dose/g) in the cell infiltrate, Spearman test. [ $^{18}\text{F}$ ]-FDG, [ $^{18}\text{F}$ ]-fluorodeoxyglucose; IL, interleukin; m-CPPD, monoclinic calcium pyrophosphate dihydrate; MSU, monosodium urate;  $\text{SUV}_{\text{max}}$ , maximum standardised uptake value.

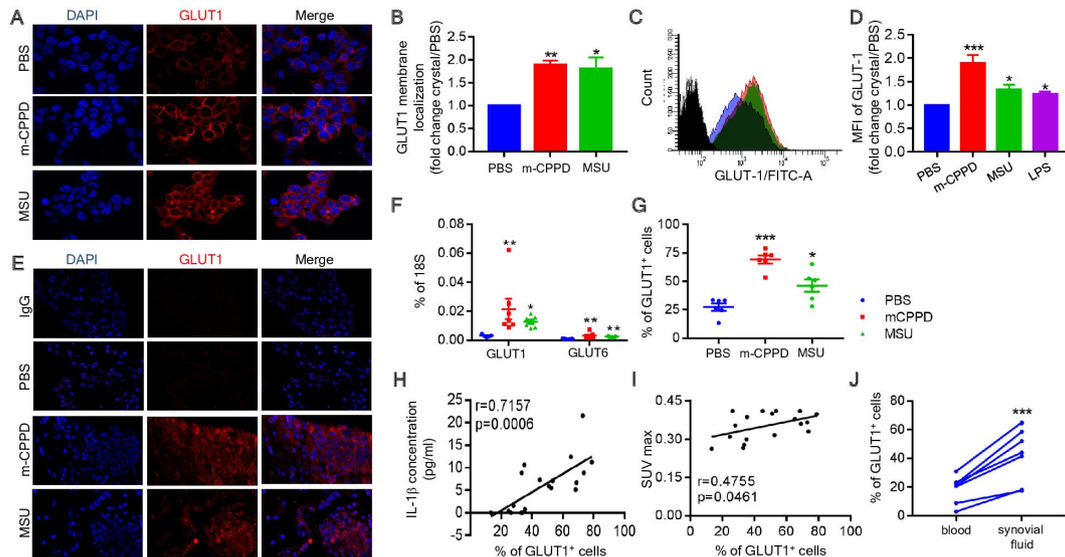
### MSU and m-CPPD crystals induce modifications in the glycolysis and TCA cycle pathways

To better understand the involvement of glucose and cell metabolism in the microcrystal inflammatory response, we performed a metabolomics study and observed in partial least squares discriminant analysis that cells stimulated by either MSU or m-CPPD crystals had robust distinct metabolic profiles without overlap components compared with unstimulated cells. Metabolomics data are available in online supplementary table S1. Specifically, we observed modifications of the abundance of multiple metabolites (figure 2A–C). Crystals induced perturbation of multiple metabolic pathways, including amino acids and glucose metabolism (see online supplementary figure S2). We observed modifications in both glycolysis pathway and TCA cycle with a slight increase of fructose-6 phosphate and a strong decrease of phosphoenolpyruvate,  $\alpha$ -ketoglutarate and malate (figure 2D,E). Beside glycolysis and TCA cycle components, there was also a decrease of glutamate and glutamine, two amino acids able to refuel the Krebs cycle in the absence of pyruvate.<sup>22 23</sup> Interestingly, the intracellular ATP was very low in crystal-stimulated cells, suggesting either an alteration of its production or an increased turnover. Then, we evaluated whether these crystals also modulated the expression of genes encoding enzymes or transporters involved in the glycolysis pathway and Krebs cycle. We did observe variations in expression of genes encoding HK-2, mitochondrial pyruvate carrier 1 and 2, monocarboxylate transporter 4, pyruvate dehydrogenase phosphatase 2, isocitrate dehydrogenase 1 and 2, and pyruvate dehydrogenase kinase 2 and 3

(see online supplementary figure S3A–D). Altogether, these results suggested that crystals altered glycolytic activity and mitochondrial function. We assessed this hypothesis with Seahorse experiments that permit analysis of real-time changes in the extracellular acidification rate (ECAR) (figure 2F) and oxygen consumption rate (OCR) (figure 2G), as surrogates of glycolysis and mitochondrial respiration, respectively. We observed that only MSU crystals increased macrophage basal glycolysis (figure 2H), glycolytic rate and glycolytic activity (figure 2I) while both MSU and m-CPPD crystals did not affect the OXPHOS (figure 2G; online supplementary figure S3E,F). Analysis of OCR suggested that MSU crystals increased only non-mitochondrial oxygen consumption (see online supplementary figure S3F).

### MSU and m-CPPD crystals increase glucose uptake associated with GLUT1 expression at the cell surface

As crystal-induced IL-1 $\beta$  production relied on glucose availability, we quantified the variation of glucose concentrations in cell culture media. By doing this, we observed that glucose concentrations were significantly lower in culture media of macrophages stimulated by either MSU or m-CPPD crystals than in culture media of unstimulated cells (figure 3A). Then, we confirmed the de novo glucose uptake by showing that MSU and m-CPPD crystals enhanced the intracellular level of radiolabelled [ $^{18}\text{F}$ ]-FDG by 152% and 148%, respectively (figure 3B). Glucose uptake has been reported in gouty joint flare in patients<sup>18 19</sup>; we reproduced this observation in the air pouch model using



**Figure 4** MSU and m-CPPD crystals increase GLUT1 expression at the cell plasma membrane. (A–C) GLUT1 membrane localisation in primed-THP-1 cells stimulated by PBS, MSU or m-CPPD was assessed by immunofluorescence confocal microscopy. (A) Imaging representative of five experiments and (B) quantification of GLUT1 membrane expression. (C) Representative overlay (MFI) by FACS of GLUT1 cell surface expression, (D) quantification of the ratio of GLUT1 MFI (crystal over PBS, LPS used as a positive control of stimulation) ( $n=5$ ). Kruskal-Wallis test with FDR correction (\*):  $*p<0.05$ ,  $**p<0.01$ ,  $***p<0.001$ . (E–I) GLUT1 expression was evaluated in mouse air pouch model of microcrystal inflammation. After PBS, MSU or m-CPPD injection (E) GLUT1 expression in the air pouch membranes was assessed by immunofluorescence confocal microscopy (representative images ( $n=5$ ), (F) expression of GLUT1 gene was assessed by RT-qPCR using RNA isolated from infiltrated cells collected in air pouch lavages. Data are represented as a % of GLUT1 or GLUT6 mRNA expression compared with 18S mRNA expression in each condition of stimulation. ( $n=6$ ). (G) Percentage of GLUT1-expressing cells was assessed by FACS using infiltrated cells collected in air pouch lavages ( $n=6$ ). Kruskal-Wallis test with FDR correction (\*):  $*p<0.05$ ,  $**p<0.01$ ,  $***p<0.001$ . Correlation between GLUT1-positive cells in the air pouch lavages and (H) IL-1 $\beta$  production or (I) glucose uptake, Spearman test. (J) Plasma membrane expression of GLUT1 was evaluated FACS comparing neutrophils isolated from flaring joint and circulating neutrophils isolated from peripheral blood of the same patient at the same moment ( $n=7$ ). Two-tailed paired t test (\*):  $*p<0.05$ ,  $**p<0.01$ ,  $***p<0.001$ . FACS, fluorescence activated cell sorting; GLUT1, glucose transporter 1; IL, interleukin; LPS, lipopolysaccharide; m-CPPD, monoclinic calcium pyrophosphate dihydrate; MFI, mean fluorescence intensity; mRNA, messenger RNA; MSU, monosodium urate; SUV<sub>max</sub>, maximum standardised uptake value.

[<sup>18</sup>F]-FDG. By PET/CT imaging, glucose uptake in response to MSU and m-CPPD crystals was evidenced by a higher labelling of the air pouch where crystals were injected, as well as higher maximum standardised uptake value (SUV<sub>max</sub>) and ratio of radio-labelled tracer found in air pouch compared with air pouch injected by saline solution (figure 3C–E). Moreover, after in vivo acquisition, quantification of the radiotracer showed that the level of [<sup>18</sup>F]-FDG was higher in air pouch membranes isolated from crystal-stimulated mice than from saline-stimulated mice (figure 3F), which was strongly correlated with IL-1 $\beta$  production in the air pouch (figure 3G).

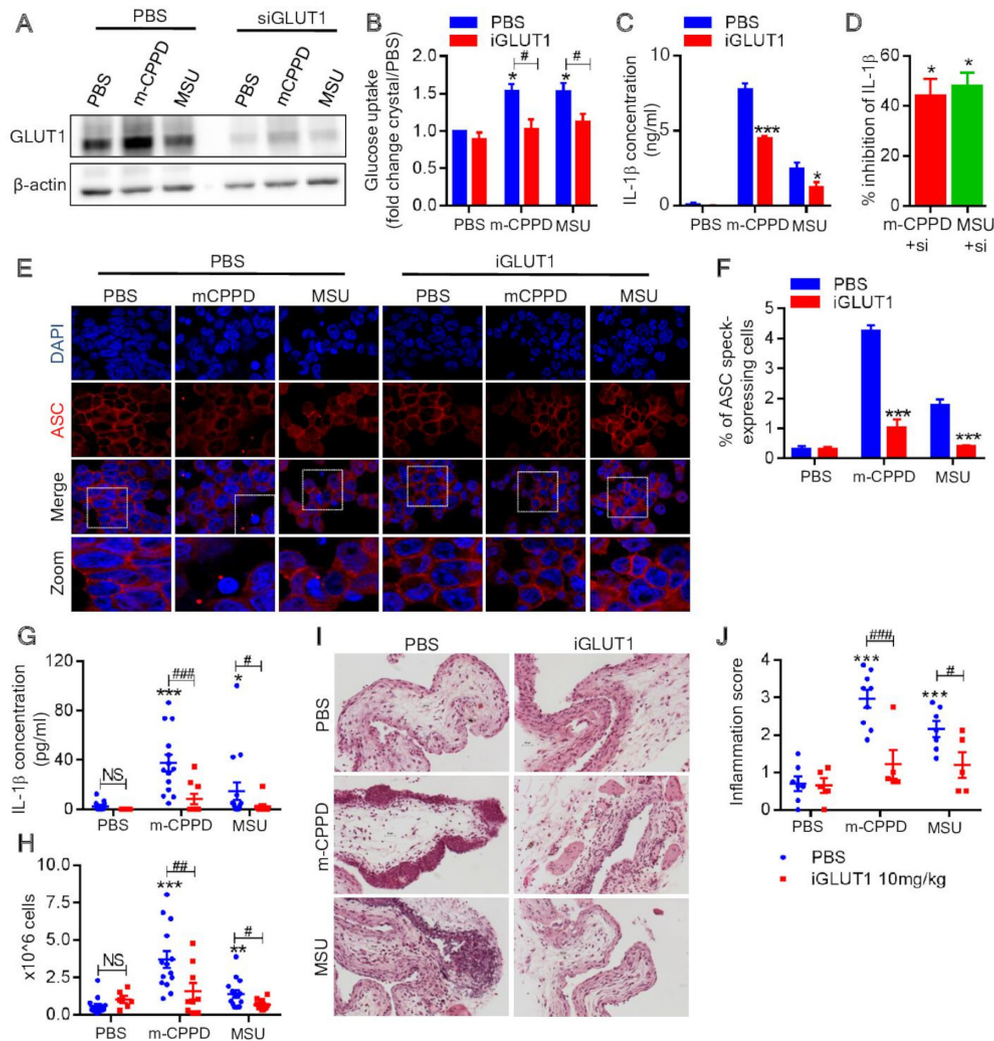
Glucose enters the cells through GLUTs that belong to solute carrier family 2 (SCL2A), which encompasses 14 members. SLC2A proteins are overexpressed in tumour cells, and SLC2A1 (GLUT1) is rapidly upregulated in inflammatory macrophages and contributes to glycolytic phenotype.<sup>24</sup> GLUT6 has been previously reported as not being involved in glucose uptake and glycolysis<sup>25</sup>; therefore, we focused on GLUT1 expression. We first showed that both MSU and m-CPPD crystal stimulation in vitro upregulated the GLUT1 gene expression on BMDMs (see online supplementary figure S4B) and the GLUT1 protein expression on THP-1 cells (figure 4A–D; see also figure 5A). By confocal microscopy, we observed that microcrystals induced a higher expression of GLUT1 at the cell plasma membrane compared with unstimulated cells (figure 4A,B). Moreover, the analysis by flow cytometry showed that both crystals, as well as the NLRP3 activator LPS, induced an increase in mean of GLUT1 expression intensity at the cell surface (figure 4C,D). The increased GLUT1 expression in response to crystals was

confirmed in vivo (figure 4E–G). First, we observed by immunofluorescence an increased expression of GLUT1 in the air pouch membrane (figure 4E). Second, the infiltrated cells into the air pouch expressed a higher level of GLUT1 messenger RNA (mRNA) (figure 4F) after crystal injection compared with saline injection, and between 50% and 70% of the recruited cells were positive for GLUT1 (figure 4G). The percentage of GLUT1-positive cells was also correlated with the inflammatory response and glucose uptake measured by IL-1 $\beta$  concentration and SUV<sub>max</sub>, respectively, in the air pouch (figure 4H,I). Finally, we validated these in vivo data by GLUT1 expression analysis on inflammatory cells isolated from gout flare patients. We observed that the ratio of GLUT1-expressing cells among the neutrophils was higher in the synovial fluid than in blood (figure 4J), suggesting that cells increased their GLUT1 expression at the site of inflammation where MSU crystals are present (MSU crystals were observed in 100% of the synovial fluid samples).

Altogether, these results suggested that MSU and m-CPPD crystal inflammation is supported by a de novo increase in GLUT1 expression and its relocalisation to the plasma membrane, required for glucose uptake and metabolic reprogramming.

### GLUT1 regulates crystal-induced inflammation

To assess the role of GLUT1 in MSU and m-CPPD crystal-induced inflammation, we adopted pharmacological (STF-31: iGLUT1) and genetic knock-down approaches (SLC2A1 small interfering RNA: siGLUT1; see validation in figure 5A) of GLUT1 inhibition. First, iGLUT1 suppressed the glucose



**Figure 5** GLUT1 drives crystal-induced inflammation. (A–F) PBS, MSU or m-CPPD crystal stimulation of primed THP-1 cells pretreated or not (PBS) with STF-31 (20  $\mu$ M, iGLUT1) or transfected with GLUT1 siRNA (si). (A) Validation of GLUT1 expression knock-down by western blot, (B) Quantification of [<sup>18</sup>F]-FDG in the cell pellets. Data shown as fold change after crystal stimulation compared with PBS. (C,D) IL-1 $\beta$  concentration measured by ELISA in the supernatants of cell culture (n=3). (D) Data are presented as % of inhibition compared with wild type THP-1 cells (n=3). (E,F) ASC speck formation was assessed by immunofluorescence confocal microscopy. (E) Imaging representative of four experiments. (F) Quantification of cells showing at least one ASC speck formation. Kruskal-Wallis test with FDR correction (\*): \*p<0.05, \*\*p<0.01, \*\*\*p<0.001. (G–J) On air pouch model of crystal inflammation in mice treated or not with GLUT1 inhibitor STF-31 (iGLUT1, 10 mg/kg). (G) IL-1 $\beta$  concentration quantified by ELISA and (H) number of infiltrate cells in air pouch lavages (n=15 per group). H&E staining (PBS: n=8 mice per group, iGLUT1: n=5 mice per group) realised on air pouch membrane sections. (I) Representative images and (J) scoring of inflammation. Two-way analysis of variance test with FDR correction (\*): \*p<0.05, \*\*p<0.01, \*\*\*p<0.001. [<sup>18</sup>F]-FDG, <sup>18</sup>F-fluorodeoxyglucose; ASC, apoptosis-associated speck-like containing a CARD; GLUT1, glucose transporter 1; IL, interleukin; m-CPPD, monoclinic calcium pyrophosphate dihydrate; MSU, monosodium urate; NS, not significant.

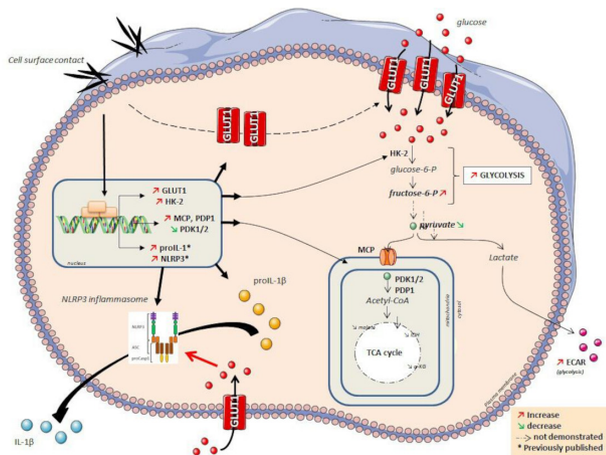
uptake induced by MSU and m-CPPD crystals (figure 5B), positioning GLUT1 as the main GLUT involved in the microcrystal response. Then, we demonstrated that GLUT1 inhibition by either of the two technical approaches led to a 50% inhibition of IL-1 $\beta$  production in response to MSU and m-CPPD (figure 5C,D). This IL-1 $\beta$  reduction under iGLUT1 was due to an inhibition of the NLRP3 inflammasome activation as GLUT1 inhibition prevented by 75% the ASC speck formation (figure 5E). Partial inhibition of NLRP3 activation and subsequently IL-1 $\beta$  production by iGLUT1 suggested that GLUT1 was involved in an amplification loop but not an initial signal of IL-1 $\beta$  production. Nevertheless, we observed that glucose depletion and GLUT1 inhibition decreased both NLRP3 and IL-1 $\beta$  gene expression and pro-IL-1 $\beta$  synthesis induced by phorbol myristate acetate (PMA) priming, suggesting that

glucose also played a role in the first signal (see online supplementary figure S5A,B).

Finally, we evaluated the iGLUT1 efficiency on microcrystal inflammation in vivo. iGLUT1 delivered by intraperitoneal injection allowed to drastically inhibit all signs of local inflammation induced by MSU and m-CPPD into the air pouch, namely, a decrease in IL-1 $\beta$  production (figure 5G), a reduction of cell infiltration (figure 5H) associated with neutrophil recruitment (see online supplementary figure S5C,D) and a global alleviation of the inflammatory score observed from HE staining of air pouch membrane (figure 5I).

## DISCUSSION

We found that MSU and m-CPPD crystal-induced macrophage production of IL-1 $\beta$ , which orchestrated the recurrent



**Figure 6** GLUT1 and glycolysis regulate NLRP3 inflammasome activation and IL-1 $\beta$  production induced by MSU and m-CPPD crystals. Crystals induced *GLUT1* gene expression, GLUT1 production and its localisation in plasma membrane. NLRP3 activation and IL-1 $\beta$  production in response to MSU and m-CPPD crystals rely on glucose uptake through GLUT1 and glycolysis. In parallel, MSU and m-CPPD crystals induce gene expression of pro-IL-1 $\beta$  and NLRP3, along with enzymes involved in glycolysis and TCA pathways, including GLUT1, HK-2, MCT, PDP1 and PDK1/2. ECAR, extracellular acidification rate; GLUT1, glucose transporter 1; HK, hexokinase; IL, interleukin; m-CPPD, monoclincal calcium pyrophosphate dihydrate; MCP, mitochondrial carrier proteins; MCT, monocarboxylate transporter; PDP, pyruvate dehydrogenase phosphatase; PDK, pyruvate dehydrogenase kinase. TCA, tricarboxylic acid.

inflammatory flares in gout and m-CPPD deposition disease, respectively, relied on glucose uptake through GLUT1 (figure 6). Glucose deprivation and glycolysis inhibition decreased NLRP3 and IL-1 $\beta$  gene expression and suppressed ASC speck formation (ie, NLRP3 activation), IL-1 $\beta$  production and neutrophil infiltration induced by MSU and m-CPPD crystals. Crystals induced GLUT1 de novo expression and its localisation to plasma membrane, along with gene expression of enzymes involved in glycolysis such as HK-2. Increase glucose uptake during MSU and m-CPPD crystal-induced inflammation has been reported in patients using [ $^{18}$ F]-FDG PET/CT.<sup>18 19</sup> We reproduced these observations in a mouse model, and we identified for the first time the key role of GLUT1 in both MSU and m-CPPD crystal-induced inflammation in mice and in human samples harvested during gout flare. Interestingly, we discovered that MSU crystal-induced glucose uptake favoured glycolytic activity, while m-CPPD crystals induced glucose uptake without modulation of glycolytic activity. Surprisingly, both microcrystals did not modulate OXPHOS. These results were distinct from the Warburg effect described in cancer cells and LPS-stimulated macrophages, where metabolic switch enhanced aerobic glycolysis at the expense of mitochondrial respiration.<sup>8-10 12 26</sup> Also reported, MSU crystals increased both OXPHOS and glycolytic activity on human neutrophils.<sup>27</sup> These findings further supported the complexity of cellular metabolic programmes and responses, which varied with the type and concentration of stimuli, cell type and cell species and tissue environment.<sup>13 28-30</sup> For example, LPS stimulated Warburg-like metabolic reprogramming in human monocytes at concentrations between 1 and 100 ng/mL but increased OXPHOS at a low dose of 0.1 ng/mL.<sup>29</sup> Likewise, monocytes stimulated with TLR2 agonist Pam3CysSK4 (P3C) or by whole microorganisms activated both glycolysis pathway and OXPHOS.<sup>29</sup> Finally, analysis of monocyte

transcriptomic responses under 28 different stimuli identified 10 clusters that represented distinct activation states with distinct metabolic responses.<sup>30</sup> These data suggested that a specific stimulus would trigger metabolic responses corresponding to a specific and functional requirement of the cells. For instance, MSU and m-CPPD crystal-induced metabolic reprogramming might increase either the phagocytosis capacity (as shown in monocytes stimulated by P3C) or the reactive oxygen species (ROS) production (as shown in neutrophils stimulated by MSU crystals),<sup>27 29</sup> two well-known mechanisms of crystal-induced NLRP3 activation.<sup>1 31</sup> In parallel, we found that MSU crystals activated macrophage glycolysis pathway to produce IL-1 $\beta$  as well described in macrophages stimulated by LPS, in activated T cells and cancer cells.<sup>8-10 12 17 26 28</sup> However, how MSU-induced and m-CPPD crystal-induced glucose uptake activates the NLRP3 inflammasome remained unknown. One explanation could come from the induction of HK-2 by MSU and m-CPPD. Indeed, in LPS-stimulated macrophages, IL-1 $\beta$  production can be modulated by metabolites and enzymes involved in the glycolysis pathway, including HK-1 and HK-2, which can both interact with mitochondrial voltage-dependent anion channel to activate NLRP3,<sup>32</sup> glyceraldehyde 3-phosphate dehydrogenase<sup>33</sup> and inactive PKM2.<sup>14 34</sup> Cancer cells and LPS-stimulated macrophages overexpress inactive dimer PKM2, which promotes HIF-1 $\alpha$  activation and transcription of HIF-1 $\alpha$ -dependent genes, including IL-1 $\beta$  and glycolytic enzymes.<sup>14 34</sup> Alternatively, MSU and m-CPPD crystal-induced glucose uptake might stimulate IL-1 $\beta$  production through Akt pathway and ROS production.<sup>27 35 36</sup> PI3K (phosphatidylinositol-3 kinase)/Akt pathway is commonly activated by MSU and m-CPPD crystals.<sup>27 36 37</sup> Akt activation enhanced IL-1 $\beta$  production through induction of ROS and glucose metabolism. Akt promoted glycolysis through stimulation of HK-2 and glucose uptake via translocation of GLUT1 to cell surface membrane and activation of its downstream mTORC1. This later stimulated HIF-1 $\alpha$ , which enhanced expression of genes involved in glycolytic reprogramming, such as *SLC2A1* (GLUT1).<sup>38</sup>

Glucose is a critical nutrient component for inflammatory macrophages, and GLUT1 is the main GLUT expressed in LPS-stimulated macrophages.<sup>24</sup> Elevated GLUT1 expression increased glucose metabolism and glycolysis, ROS production and expression of proinflammatory mediators, including IL-1 $\beta$ .<sup>24</sup> Here, we found that MSU and m-CPPD crystals triggered plasma membrane localisation of GLUT1, which drove glucose uptake, NLRP3 activation and IL-1 $\beta$  production. Furthermore, we observed in gouty patients that GLUT1 was more frequently expressed at the surface of neutrophils isolated from flaring joint than neutrophils isolated from peripheral blood of the same patient. These results added crystal-induced IL-1 $\beta$  production to the list of inflammatory conditions regulated by GLUT1-induced glucose uptake in macrophages, such as cancer, infection, autoimmune disease, diabetes and obesity.<sup>14 24 28 35 39 40</sup> Interestingly GLUT1 also governed post-prandial glucose uptake by peritoneal macrophages, leading to IL-1 $\beta$  production and, subsequently, insulin secretion.<sup>35</sup> This might explain why gout flare frequently occurred after a feast. MSU and m-CPPD crystals enhanced both GLUT1 mRNA expression and GLUT1 plasma membrane localisation by unknown mechanisms. GLUT1 activity is regulated by its membrane localisation and activation, kinetic of its internalisation, endosomal sorting and recycling back to the cell membrane, which depends on retromer cargo complex.<sup>41 42</sup> Cancer cells favour aerobic glycolysis by promoting GLUT1 plasma membrane localisation through the PI3K/Akt/mTORC

pathway.<sup>43–45</sup> Akt activation promotes cell surface membrane recycling of GLUT1 and reduces its internalisation.<sup>43</sup> Inversely, Akt inhibition by phosphatase TENSin homologue deleted on chromosome 10 (PTEN) prevented GLUT1 plasma membrane localisation.<sup>45–47</sup> Whether MSU and m-CPPD crystals regulated GLUT1 membrane expression through the Akt pathway needs to be studied. Alternatively, MSU and m-CPPD crystals might induce GLUT1 membrane localisation through thioredoxin-interacting protein, which facilitated GLUT1 endocytosis via clathrin-coated pits, modulated PTEN activity and was involved in crystal-induced inflammation.<sup>31, 48</sup> Lastly, crystal-induced glucose uptake might be due to an increase in GLUT1 transport activity. Indeed, the drastic decrease of cytosolic ATP observed after MSU and m-CPPD crystal stimulation might stimulate GLUT1 transport activity, as previously reported.<sup>49, 50</sup>

Although *SLC2A6* (GLUT6) gene expression in THP-1 cells was highly enhanced by either MSU or m-CPPD crystal stimulation, its exact role in crystal-induced inflammation and glucose uptake is currently unknown. Recent data suggested that GLUT6 was involved in neither glucose uptake nor glycolysis nor OXPHOS.<sup>25, 51, 52</sup> GLUT6-deficient mice had normal glucose metabolism<sup>51</sup> and GLUT6-deficient BMDMs had similar ECAR and OCR than wild-type cells.<sup>25, 52</sup> Moreover, expression of proinflammatory mediators such as IL-1 $\beta$  and TNF- $\alpha$  was unchanged in the absence of GLUT6.<sup>52</sup> Further studies are needed to understand how GLUT6 participates to crystal-induced inflammation.

How MSU and m-CPPD crystals induced GLUT1 membrane localisation constituted the main limitation of our study. Although we did not identify the exact mechanisms involved in crystal effects, our results suggested that MSU and m-CPPD crystals increased, in THP-1 and mouse BMDM cells, GLUT1 gene expression, GLUT1 protein production and membrane location and GLUT1 activity. Whether crystals induced IL-1 $\beta$  production by primary human monocytes does also depend on glucose uptake need specific studies. Similarly, we did not assess in this study how hyperglycaemic modulated crystal-induced inflammation in vivo. We planned to address this question in type 2 and type 1 diabetes using Ob/Ob mice and streptozotocin mouse models, respectively. Interestingly, recent report supported our findings by showing that patients treated with metformin had less gout flare.<sup>53</sup> Finally, metabolic reprogramming during gout flare needed also to be confirmed in patients by doing metabolomics analysis in neutrophils isolated from inflamed joint.

Overall, metabolic changes characterised by GLUT1-mediated glucose uptake and increase in glycolysis governed MSU and m-CPPD crystal-activated IL-1 $\beta$  production. These findings open new therapeutic paths to modulate crystal-related inflammation.

## MATERIALS AND METHODS

See a fully detailed Materials and methods in the online supplementary file.

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**Acknowledgements** We thank Mylène Zarka Prost-Dumont, Morgane Bourmaud, Yohan Jouan and Yetki Aslan (UMRS-1132) for their precious help during the in vivo experiments.

**Contributors** H-KE conceived the study. FR, LC-G and H-KE contributed to its design and coordination, participated in data interpretation and cowrote the manuscript. LC-G and FR performed the laboratory experiments. CC synthesised calcium pyrophosphate crystals, characterised their physicochemical structure and contributed to writing the manuscript. LO, FA and NV performed the Seahorse experiments. FC and FF realised the metabolomics study. AP performed PET study. PR, TB, FR and H-KE collected the patient samples. H-KE, CC, PR, MC-S and FL secured the funding. All authors participated in the final approval of the manuscript.

**Funding** The study was funded by ANR (ANR-126BS08-0022-01), ART Viggo, the 'Prevention et Traitement des Décalsifications (PTD)' Association, Arthritis Courtin foundation (Arthritis R&D 2018-2019) and the French Society of Rheumatology (SFR 2017-2018, SFR 2018-2019). LC-G was financially supported by grants from ANR and ART Viggo, and FR by Paris Diderot University and ART Viggo, PTD.

**Competing interests** None declared.

**Patient and public involvement** Patients and/or the public were not involved in the design, conduct, reporting or dissemination plans of this research.

**Patient consent for publication** Not required.

**Provenance and peer review** Not commissioned; externally peer reviewed.

**Data availability statement** No data are available. All data relevant to the study are included in the article or uploaded as supplementary information. All data are included in the article.

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