CDK8 Fine-Tunes IL-6 Transcriptional Activities by Limiting STAT3 Resident Time at the Gene Loci

Highlights
- CDK8 regulates IL-6-mediated STAT3 S727 phosphorylation in primary human T cells
- CDK8 controls STAT3 activity by limiting its resident time at gene loci
- CDK8 inhibition increases IL-6-mediated Th17 differentiation

Authors
Jonathan Martinez-Fabregas, Luopin Wang, Elizabeth Pohler, ..., Majid Kazemian, Suman Mitra, Ignacio Moraga

Correspondence
kazemian@purdue.edu (M.K.), suman.mitra@inserm.fr (S.M.), imoragagonzalez@dundee.ac.uk (I.M.)

In Brief
How IL-6 elicits its immune pleiotropic activities is not fully understood. Martinez-Fabregas et al. show that CDK8 represses IL-6-mediated transcription by limiting STAT3 resident time at the gene loci. By regulating CDK8 expression levels, immune cells can adapt their responses to STAT3-activating cytokines.
CDK8 Fine-Tunes IL-6 Transcriptional Activities by Limiting STAT3 Resident Time at the Gene Loci

Jonathan Martinez-Fabregas,1,6 Luopin Wang,2,6 Elizabeth Pohler,1 Adeline Cozzani,4 Stephan Wilmes,1 Majid Kazemian,2,5,* Suman Mitra,4,5,* and Ignacio Moraga1,5,7,*

1Division of Cell Signaling and Immunology, School of Life Sciences, University of Dundee, Dundee, UK
2Department of Computer Science, Purdue University, West Lafayette, IN, USA
3Department of Biochemistry, Purdue University, West Lafayette, IN, USA
4Université de Lille, INSERM UMR1277 CNRS UMR9020–CANTHER and Institut pour la Recherche sur le Cancer de Lille (IRCL), Lille, France
5Senior author
6These authors contributed equally
7Lead Contact
*Correspondence: kazemian@purdue.edu (M.K.), suman.mitra@inserm.fr (S.M.), imoragagonzalez@dundee.ac.uk (I.M.)

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SUMMARY

Cytokines are highly pleiotropic ligands that regulate the immune response. Here, using interleukin-6 (IL-6) as a model system, we perform detailed phosphoproteomic and transcriptomic studies in human CD4+ T helper 1 (Th-1) cells to address the molecular bases defining cytokine functional pleiotropy. We identify CDK8 as a negative regulator of STAT3 transcriptional activities, which interacts with STAT3 upon IL-6 stimulation. Inhibition of CDK8 activity, using specific small molecule inhibitors, reduces the IL-6-induced phosphoproteome by 23% in Th-1 cells, including STAT3 S727 phosphorylation. STAT3 binding to target DNA sites in the genome is increased upon CDK8 inhibition, which results in a concomitant increase in STAT3-mediated transcriptional activity. Importantly, inhibition of CDK8 activity under Th-17 polarizing conditions results in an enhancement of Th-17 differentiation. Our results support a model where CDK8 regulates STAT3 transcriptional processivity by modulation of its gene loci resident time, critically contributing to diversification of IL-6 responses.

INTRODUCTION

Cytokines are critical orchestrators of innate and adaptive immunity (Lin and Leonard, 2019). Despite the functional relevancy of this family of ligands, the molecular basis governing their large functional pleiotropy remains poorly defined. Cytokines exert their activities by dimerizing/oligomerizing surface receptors and triggering the tyrosine (Tyr) phosphorylation of STAT transcription factors by janus kinases (JAKs) (Gorby et al., 2018; Martinez-Fabregas et al., 2019; Stroud and Wells, 2004; Wang et al., 2009; Wilmes et al., 2020). This in turn leads to the nuclear translocation of STATs and the induction of specific gene expression programs and bioactivities (Poli and Camporeale, 2015; Schindler et al., 2007). However, how qualitative and quantitative changes in these pathways contribute to cytokine functional pleiotropy is poorly understood.

STATs can be modified in conserved Tyr or serine (Ser) residues (Decker and Kovarik, 2000). Although STAT Tyr phosphorylation plays a critical role in mediating cytokine responses, the role of STAT Ser phosphorylation in cytokine-mediated activities is less clear (Chung et al., 1997; Decker and Kovarik, 2000; Kim and Baumann, 1997; Wen et al., 1995). Early work in cancer cell lines showed that Ser phosphorylation of STAT proteins regulated their transcriptional activities (Wen et al., 1995). However, whether STAT Ser phosphorylation promotes a positive or negative effect on STAT transcriptional activities remains more controversial (Bancerek et al., 2013; Decker and Kovarik, 2000; Levy and Darnell, 2002; Lim and Cao, 1999). Some studies report a positive effect of STAT Ser phosphorylation in driving STAT transcriptional activities, whereas others have reported an opposite effect (Bancerek et al., 2013; Decker and Kovarik, 2000; Levy and Darnell, 2002; Lim and Cao, 1999; Steen et al., 2016; Wen et al., 1995; Yokogami et al., 2000). For STAT3, Ser phosphorylation appears to negatively impact its transcriptional activities by regulating its chromatin binding dwell time (Yang et al., 2020). But, earlier studies had reported that STAT3 Ser phosphorylation did not have effects on STAT3 chromatin binding (Wen and Darnell, 1997). The differences in these observations could be attributed to the use of different cancer cell lines, suggesting a context-dependent regulation of STAT3 signaling.

Interleukin-6 (IL-6) represents a classical paradigm for cytokine functional pleiotropy. IL-6 acts as a central regulator of the immune response by triggering both pro-inflammatory and anti-inflammatory responses (Hunter and Jones, 2015; O’Shea and Murray, 2008; Rose-John, 2018; Scheller et al., 2011). IL-6 drives inflammatory processes by modulating the adaptive and innate immunity arms. On the one hand, IL-6 promotes the differentiation of T helper-17 (Th-17) cells, while inhibiting the
differentiation of T regulatory (T reg) cells (Jones et al., 2010; Korn et al., 2008). On the other hand, IL-6 recruits myeloid cells to sites of inflammation (Fielding et al., 2008; Gabay, 2006). Additionally, dysregulation of IL-6 or IL-6-mediated responses is often associated with inflammatory disorders, making this cytokine highly relevant for human health (Jones and Jenkins, 2018; Tanaka et al., 2014). IL-6 exerts its activities by triggering the activation of the JAK1/STAT1/STAT3 signaling pathway upon recruitment of IL-6Rα and gp130 receptor subunits (Heinrich et al., 1998; Martinez-Fabregas et al., 2019; Servais et al., 2019). However, despite the critical contribution that IL-6 plays in modulating the immune response, the role that STAT3 Ser phosphorylation plays on regulating IL-6 immune activities has not been explored in detail. Studies on STAT1 have shown that CDK8 appears to be the kinase driving its Ser phosphorylation in response to interferon gamma (IFNγ) stimulation in macrophages (Bancerek et al., 2013). Blockage of STAT1 Ser phosphorylation, using STAT1 Ser mutants, significantly altered the IFNγ transcriptional response and its ability to clear Listeria monocytogenes infection, highlighting the relevance of STAT1 Ser phosphorylation in modulating IFNγ response in macrophages (Varinou et al., 2003). However, how Ser phosphorylation fine-tunes STAT3 immuno-modulatory activities is less well known.

In this study, we set out to characterize how the signaling initiated by IL-6 in human T cells leads to its functional pleiotropy. We detected both STAT3 Tyr and Ser phosphorylations in response to IL-6 stimulation, with STAT3 Ser phosphorylation exhibiting a delayed activation. Using a battery of small molecule inhibitors, we identified CDK8 and CDK9 as the kinases driving Ser phosphorylation of STAT3 upon IL-6 stimulation in T cells. Using proximity ligation studies, we confirmed the increased interaction between STAT3 and CDK8 and CDK9 in the nucleus upon IL-6 stimulation. Inhibition of the activity of these two kinases resulted in a more robust interaction with STAT3, even in the absence of IL-6 stimulation. Chromatin immunoprecipitation sequencing (ChIP-seq) and RNA sequencing (RNA-seq) studies revealed a global increase in STAT3 chromatin binding upon inhibition of CDK8, resulting in the induction of a larger gene expression program by IL-6. In agreement with this enhanced gene expression program, IL-6 induced a more robust differentiation of Th-17 cells upon CDK8 inhibition in vitro. Overall, our studies identify a STAT3 regulatory mechanism in T cells, whereby CDK8 and CDK9 modulate STAT3 processivity by controlling its chromatin binding dwell time and transcriptional activity. These observations suggest ways to manipulate IL-6- and STAT3-mediated responses by fine-tuning CDK8/9 activities.

**RESULTS**

**IL-6 Signaling Preferentially Induces Tyr/Ser Phosphorylation of STAT1/3 in Human T Cells**

IL-6 critically contributes to modulating the T cell response. Yet, we have a poor understanding of the signaling networks engaged by IL-6 in T cells and their role in shaping IL-6 immune activities. To gain insight into the IL-6 signaling network in T cells, we carried out detailed signaling studies in human resting or activated CD4+ and CD8+ T cells stimulated with IL-6. IL-6 receptor expression varies significantly among different T cell populations and environmental contexts (Figure S1; Betz and Müller, 1998; Jones et al., 2010; Ridgley et al., 2019), making the study of IL-6 signaling in T cells challenging. To minimize this variability, we have used Hyper-IL-6 (HyIL-6) for our signaling studies. HyIL-6 is a synthetic heterodimer comprised of IL-6Rα and IL-6 proteins connected by a gly/ser linker (Fischer et al., 1997). HyIL-6 triggers signaling in all cells expressing gp130, producing a more robust and homogeneous signaling output (Rose-John, 2012). In Th-1 cells, HyIL-6 induced a more potent phosphorylation of STAT1 and STAT3 than IL-6 (Figure S1A, left panels). However, the pSTAT1/pSTAT3 ratio induced by the two ligands was identical (Figure S1A, right panel), suggesting that HyIL-6 exhibits only qualitative and no quantitative signaling differences with IL-6. Dose-response (Figure 1A) and kinetic signaling studies (Figure 1B) showed that resting and activated CD4+ and CD8+ T cells respond to HyIL-6 treatment by Tyr phosphorylating STAT1 and STAT3 transcription factors. The STAT1/3 activation amplitudes elicited by HyIL-6 in these cells, however, differ significantly, with activated CD4+ and CD8+ T cells triggering between 2- to 3-fold higher STAT1/3 phosphorylation amplitudes than resting CD4+/CD8+ T cells in response to HyIL-6 treatment (Figures 1A and 1B). Interestingly, activated CD4+ T cells triggered higher levels of STAT1 phosphorylation than activated CD8+ T cells upon HyIL-6 stimulation (Figures 1A and 1B, left panels), which correlated with a higher expression of gp130 and IL-6Rα by CD4+ T cells (Figure S1B). Overall, these results show that IL-6-induced signaling in human T cells is a dynamic and context-dependent process.

To gain further insight into the signaling networks, beyond JAK/STAT1/3, engaged by IL-6 in T cells, we used an antibody array targeting 28 relevant signaling intermediaries. Resting and activated CD4+ and CD8+ T cells were stimulated with saturating concentrations of HyIL-6 for the indicated times, and their signaling signatures in response to HyIL-6 treatment were assayed by flow cytometry (Figures 1C and 1D). To ensure the quality of our signaling antibody array, we stimulated different populations of T cells with anti-CD3/anti-CD28 antibodies (TCR)+IL-2 as a positive control, because this treatment activates a large proportion of the signaling molecules detected by our antibody array (Ross et al., 2016; Smith-Garvin et al., 2009). In both resting and activated human CD4+ and CD8+ T cells, TCR+IL-2 treatment led to the activation of a large proportion of the signaling intermediaries, including STAT1, STAT3, STAT4, STAT5, STAT6, ERK, AKT, S6R, and CREB (Figures 1C and 1D; Figures S2 and S3). HyIL-6 treatment on the other hand preferentially induced the Tyr and Ser phosphorylation of both STAT1 and STAT3 and, to a lower extent, STAT4 (Figures 1C and 1D; Figure S2). STAT1 and STAT3 Tyr and Ser phosphorylation in response to HyIL-6 treatment were inhibited by the JAK inhibitor tofacitinib, confirming the dependency of these two modifications on JAK activity (Figures 1E and 1F; Figures S3A–S3D). Overall, our signaling data support a dynamic activation of the JAK/STAT pathway by IL-6 in T cells, suggesting that IL-6 functional pleiotropy emanates from activation of a few signaling intermediaries.
IL-6 Induces a Large Number of Phosphoproteome Changes in Human T Cells

To obtain a full spectrum of the IL-6 signalosome in T cells, we next performed a quantitative high-resolution phosphoproteomics assay using stable isotope labeling by amino acids in cell (SILAC). We selected Th-1 cells due to their significance in immunity and ability to expand in vitro to large quantities of a highly pure population, which is compatible with SILAC studies (Figure 2A).

Figure 1. IL-6 Signaling Landscape in Primary Human T Cells

(A and B) STAT1 and STAT3 phosphorylation in response to various doses (A) and exposure time (B) of IL-6 stimulation in resting and activated primary human CD4+ and CD8+ T cells. Error bars show mean ± SEM from three individual biological replicas.

(C and D) Phospho-FLOW analysis of IL-6 signaling pathways in resting (C) and activated primary human CD4+ and CD8+ T cells treated with HyLL-6 or anti-CD3/CD28 (TCR) + IL-2. ns, cells without any stimulation. Heatmaps show fold change in the level of phosphorylation or protein expression of the different proteins. See also Figures S2 and S3.

(E and F) Effect of JAK inhibition (2 μM tofacitinib) on the phosphorylation of STAT1 (E) and STAT3 (F) Tyr701 and Ser727 in resting and activated primary human CD4+ and CD8+ T cells. Error bars show mean ± SEM from three individual biological replicas.
Figure 2. Phosphoproteomic Landscape of IL-6 in Human Primary CD4+ Th-1 Cells
(A) Experimental workflow for SILAC-based quantitative phosphoproteomic analysis of human primary CD4+ Th-1 cells stimulated with 20 nM HyIL-6 for 15 min.
(B) Volcano plot showing differential phosphopeptides in unstimulated versus stimulated Th-1 cells with 20 nM HyIL-6 for 15 min. Phosphopeptides identified in six biological replicates are shown as log-transformed SILAC ratios plotted against log-transformed p values (two-sided t test). Phosphosites changed more than 1.5-fold with a p value of <0.05 are shown in red (decreased) or blue (increased). Select phosphopeptides are labeled (see also Table S1 for full list). The 24 phosphosites more reproducibly decreased (red) or increased (blue) are displayed alongside.
The combined analysis from six independent biological replicates of unstimulated versus HyIL-6-stimulated Th-1 cells identified 17,935 phosphosites on 4,196 proteins (Figure 2B; Table S1). Among those phosphosites, 304 were increased and 245 were decreased significantly in response to HyIL-6 stimulation in human primary Th-1 cells, while the rest remained unchanged (Figure 2B). Gene Ontology (GO) analysis (KEGG, Kyoto Encyclopedia of Genes and Genomes) pathways and molecular function analyses showed an enrichment of the JAK/STAT pathway, as expected, upon stimulation of Th-1 cells with HyIL-6 (Figure 2C). Of all proteins with detected phosphosites, 288 were kinases (Figure 2D; Table S1). Of those proteins, 28 were regulated by HyIL-6 treatment (Figure 2D; Table S1) and were enriched in several signaling pathways (e.g., mTOR and mitogen-activated protein kinase [MAPK]) (Figure 2E). GO analysis showed that 41% of the phosphoproteomic changes induced by HyIL-6 took place in the nucleus (Figures S3B and S4), highlighting this compartment as an important signaling platform for IL-6 activities. Furthermore, our GO analysis indicated an enrichment of phosphosites involved in the regulation of transcription and more specifically RNA polymerase II (RNA Pol II)-mediated transcription (Figure 2C). HyIL-6 treatment regulated processes related to protein transcription, acting as part of DNA-modifying complexes (MTA1, CCAR2, and TRRAP) or serving as transcriptional regulators (ELF2, RUNX2, TSC22D4, and SP4); histone (de)acetylases (KANSL2, TRRAP, and RBBP7); RNA Pol II transcription (MECPE, MEF2C, and MED1); and mRNA splicing (DDX46 and SF3B4), processing (RBM6 and RBM39), and export (NUP50 and NUP153) (Figures S3B and S4). HyIL-6 treatment also regulated non-nuclear processes, including regulation of the cytoskeleton, translation, and proteasome at the cytoplasmic level (Figure 2C; Figures S3 and S4). Overall, our phosphoproteomic data revealed a strong regulation of the nuclear phosphoproteome by IL-6 that could contribute to fine-tuning its immunomodulatory activities.

**CDK8/CDK9 Regulate STAT1 and STAT3 Ser727 Phosphorylation**

HyIL-6 triggers the Tyr and Ser phosphorylation of STAT1 and STAT3 in T cells. Although JAK1 contributes to the Tyr phosphorylation of STAT1/3, the kinase responsible for STAT1/3 Ser phosphorylation in T cells is currently not known. To identify this kinase, we used a panel of inhibitors targeting signaling pathways previously shown to regulate STAT1 and STAT3 Ser phosphorylation in different cellular systems (Decker and Kovarik, 2000). Tofacitinib, a JAK inhibitor, blocked both STAT1/STAT3 Tyr and Ser phosphorylation by HyIL-6 treatment (Figures 3A and 3B), confirming previous observations. Of the battery of inhibitors tested, only Torin 1, an mTOR inhibitor targeting both mTORC1 and mTORC2 complexes (Thoreen et al., 2009), inhibited the Ser phosphorylation induced by HyIL-6 in both STAT1 and STAT3, without affecting their Tyr phosphorylation (Figures 3A and 3B). Rapamycin, an inhibitor that under our experimental conditions only targets the mTORC1 complex (Laplanche and Sabatini, 2012; Sarbassov et al., 2006), failed to do so, suggesting that the Ser phosphorylation of STAT1 and STAT3 induced by HyIL-6 was a mTORC2-mediated response (Figures 3A and 3B). However, alternative mTOR inhibitors (i.e., AZD8055 and KU0063794) failed to restrict STAT1 and STAT3 Ser phosphorylation by HyIL-6, indicating that the Torin1-mediated inhibition was an off-target effect (Figure 3C). Moreover, inhibitors specifically targeting well-described off-targets of Torin 1, i.e., ataxia telangiectasia mutated (ATM; KU55933) and DNA-dependent protein kinase (DNA-PK; KU57788) (Liu et al., 2012), failed to inhibit STAT1 and STAT3 Ser phosphorylation by HyIL-6 (Figure 3D). This finding indicates that the inhibition of STAT1 and STAT3 Ser phosphorylation is a previously unknown off-target effect of Torin1 and needs to be accounted for when used in in vitro or in vivo studies.

Previous studies have described CDKs as regulators of STAT1 Ser phosphorylation in different systems (Bancerek et al., 2013; Chen et al., 2019; Kosciuczuk et al., 2019; Putz et al., 2013). Thus, we next tested whether the STAT3 Ser phosphorylation induced by HyIL-6 was mediated by CDKs. For that test, we measured STAT3 Tyr and Ser phosphorylation levels induced by HyIL-6 in cells treated with a panel of CDK inhibitors. Flavopiridol, a well-described pan-CDK inhibitor (Luke et al., 2012), completely abolished the Ser phosphorylation of STAT3 induced by HyIL-6 (Figure 3E; Figures S5A–S5C). CDK8-specific inhibitors (i.e., BI-1347 in Hofmann et al., 2020 and MSC2530818) inhibited HyIL-6-induced STAT3 Ser phosphorylation by 70% in both human CD4+ and CD8+ T cells (Figure 3E; Figures S5A–S5C). CDK9 (i.e., NVP2 and CDK inhibitor II) or CDK12/CDK13 (i.e., THZ531 and MFH-2-90-1) inhibitors only reduced the STAT3 Ser phosphorylation by 20% (Figure 3E; Figures S5A–S5C). None of the inhibitors affected the STAT3 Tyr phosphorylation levels (Figure 3E, Figures S5A–S5C). Genetic silencing of CDK8 and CDK9 in HEK293T cells (Figures S5D–S5F) further support CDK8 and CDK9 as the main kinases driving STAT3 Ser phosphorylation in response to HyIL-6. We detected a small decrease on STAT3 Tyr phosphorylation upon HyIL-6 stimulation in cells depleted of CDK8 and CDK9 (Figure S5E), which we did not detect in experiments using small molecule inhibitors. We believe that this decrease results from toxicity associated with prolonged depletion of CDK8 and CDK9 kinases. Our data highlight a critical role of CDK8 in regulating STAT3 Ser phosphorylation by HyIL-6, with an accessory role of other CDK members.

**HyIL-6 Induces Nuclear Interaction of STAT3 and CDK8/CDK9**

Next, we explored whether STAT3 and CDK8/9 physically interacted in the nucleus upon HyIL-6 stimulation. For that, we performed proximity ligation assays (PLAs), a technique that allows the detection of protein complexes at endogenous levels without the need of protein overexpression or labeling that could interfere with their binding partners (Fredriksson et al., 2002).
Activated human primary CD4+ T cells were stimulated with 20 nM HyIL-6 for the indicated times, and samples were prepared for PLA analysis following manufacturer instructions (Sigma). In untreated cells, we detected very low levels of STAT3/CDK8 (Figure 4A) and STAT3/CDK9 complexes (Figure 4B). Upon HyIL-6 stimulation, we detected a 2- to 4-fold increase in the number of STAT3/CDK8 and STAT3/CDK9 complexes, which peak at 30 min after stimulation and return to basal levels after 3 h (Figures 4A and 4B), paralleling the STAT3 Tyr activation kinetics (Figure 1B).

We next studied whether CDK activity modulated the formation of STAT3/CDK complexes upon HyIL-6 stimulation. Activated human CD4+ T cells were stimulated with HyIL-6 for 30 min in the presence of different CDKs inhibitors. Levels of STAT3/CDK8 and STAT3/CDK9 complexes were measured by PLA analysis. As before, HyIL-6 stimulation led to a significant increase in the number of STAT3/CDK8 and STAT3/CDK9 complexes, when compared to unstimulated cells (Figures 4C and 4D). Addition of flavopiridol (panCDK inh.) or MSC2530818 (CDK8 inh.) inhibitors resulted in an enhancement in the number of STAT3/CDK8 and STAT3/CDK9 complexes (Figures 4C and 4D). Interestingly, treatment with the two inhibitors led to an increase in the number of STAT3/CDK8 and STAT3/CDK9 complexes (Figures 4C and 4D) in the absence of HyIL-6 stimulation, suggesting a role for CDK activities in regulating STAT3 nuclear resident time and thus chromatin binding.

Previous studies have reported that mutation of S727 in STAT3 to alanine (Ala) modulates its transcriptional activity (Chung et al., 1997; Kim and Baumann, 1997; Lim and Cao, 1999; Wen et al., 2006).
A

HyL6 (min) 0 5 30 60 180

DAPI

CDK8/STAT3

B

HyL6 (min) 0 5 30 60 180

DAPI

CDK8/STAT3

C

HyL6 20nM MSC2530818 Flavopiridol

DAPI

CDK8/STAT3

D

HyL6 20nM MSC2530818 Flavopiridol

DAPI

CDK9/STAT3

E

HyL6 (min) 0 5 30 60 180

DAPI

STAT3 WT - GFP

CDK9/STAT3

DAPI

STAT3 S727A

CDK9/STAT3

(legend on next page)
Thus, we next investigated the role that this mutation plays on recruitment of CDKs upon HyIL-6 stimulation. For that investigation, we took advantage of the human Hut78 cell line, a cutaneous T lymphocyte, where HyIL-6 treatment also induced STAT3 S727 phosphorylation in a CDK-dependent manner (Figure S6A-B). Importantly, only flavopiridol treatment resulted in an inhibition of STAT3 Ser phosphorylation by HyIL-6 in Hut78 cells, suggesting that in these cells, CDK9 but not CDK8 is the main CDK driving STAT3 phosphorylation (Figures S6A and S6B). Next, we generated STAT3 knockdown (STAT3 KnD) Hut78 cell lines by CRISPR-Cas9 (Figure S6C). These cells exhibited a clear reduction in the STAT3 Tyr phosphorylation upon HyIL-6 stimulation (Figure S6D). STAT3 KnD cells were reconstituted with wildtype (WT) STAT3-GFP or S727A STAT3-GFP mutant, and the levels of STAT3/CDK9 complex formation were measured by PLA (Figure 4E; Figure S6E). Due to STAT3 overexpression in these cells, we detected significantly higher levels of the STAT3/CDK9 complex in unstimulated cells than those detected in human Th-1 cells. Yet, upon HyIL-6 stimulation, we observed a significant increase in the number of STAT3 WT/CDK9 complexes in the nucleus, which peaked at 30 min and went back to basal levels by 2 h after treatment (Figure 4E). The STAT3 S727A mutant exhibited a similar nuclear translocation profile to STAT3 WT, but it showed a delayed association kinetic with CDK9 (Figure 4E). Interestingly, at late stimulation times, we observed higher levels of STAT3 S727A/CDK9 complexes when compared to STAT3 WT, suggesting that the STAT3/CDK9 interaction is stabilized in the absence of STAT3 Ser phosphorylation (Figure 4E). Overall, our studies show that CDK8 and CDK9 fine-tune STAT3 nuclear dynamics.

CDK8 Regulates IL-6-Induced Nuclear Phosphoproteome

Our data have highlighted a critical role of CDK8 in regulating STAT3 Ser phosphorylation and nuclear dynamics in human Th-1 cells (Figures 3E, 4A, and 4C). Next, we asked which proportion of the IL-6-regulated phosphoproteome was dependent on CDK8 activity. For that investigation, we performed phosphoproteomics studies in Th-1 cells stimulated with HyIL-6 for 15 min in the absence or presence of the CDK8 inhibitor MSC2530818. The combined analysis of our phosphoproteomics study identified 11,035 phosphosites in 3,500 proteins (Figures 5A and 5B; Table S2). To minimize mis-interpretation of the data resulting from off-target effects mediated by the CDK8 inhibitor, we focused our analysis on phosphosites that were induced by HyIL-6 treatment and sensitive to CDK8 inhibition. HyIL-6 treatment induced 162 and repressed 160 phosphosites, of which 88 of them (63 of the upregulated and 25 of the downregulated) were sensitive to CDK8 inhibition (Figures 5A and 5B; Table S2). Consistent with our initial phosphoproteome study (Figure 2), a large fraction (34%) of the phosphoproteomic changes induced by HyIL-6 took place in the nuclei of the cells, based on GO analysis (Figure 5C). GO analysis studies indicated that HyIL-6 stimulation regulated proteins involved in transcription, specifically RNA-Pol-II-mediated transcription and other cellular processes such as histone (de)acetylation and DNA methylation (Figures 5D and 5E). A total of 27% of proteins involved in transcription and 40% of proteins involved in RNA Pol II transcription were affected by CDK8 inhibition (Figure 5F).

A schematic view of the nuclear IL-6-induced phosphoproteome and its regulation by CDK8 is presented in Figure 5G. Overall, our data highlight a critical contribution of CDK8 in shaping the IL-6 phosphoproteome by regulating processes associated with RNA-Pol-II-mediated transcription.

CDK8 Regulates STAT3-Mediated Transcription

We showed that STAT3 nuclear dynamics were regulated by CDK8 activity. Thus, we next asked whether inhibition of CDK8 activity would alter STAT3-dependent gene transcription. For that question, we performed RNA-seq studies in Th-1 cells stimulated with HyIL-6 with or without the CDK8 inhibitor MSC2530818 for 6 h (Figures 6A–6D; Figure S7A). CDK8 inhibition by MSC2530818 neither altered the phosphorylation profile of RBP1 nor blocked transcriptional upregulation induced by HyIL-6 treatment, suggesting that CDK8 inhibition did not result in an overall transcription blockage (Figure 6; Figure S7B) (Czodrowski et al., 2016; Harlen and Churchman, 2017). As previously observed by our laboratory (Martinez-Fabregas et al., 2019), HyIL-6 stimulation alone resulted in changes in the expression pattern of a small number of genes (n = 27) including classical STAT3 targets in Th-1 cells (Figures 6A and 6B, left panel). Treatment with only CDK8 inhibitor led to changes in the expression of 111 genes, of which 84 were upregulated (Figures 6A and 6B, middle panel). The combined HyIL-6 and MSC2530818 treatments exhibited a synergistic effect, leading to changes in the expression of 176 genes (Figures 6A and 6B, right panel), suggesting that CDK8 inhibition induced transcriptional programs by promoting HyIL-6/STAT3-mediated transcription. Differently regulated genes seem to fall into a few categories: genes induced by HyIL-6 treatment, but not regulated by CDK8 inhibition (e.g., BCL3 and SOCS3); genes induced...
up upon CDK8 inhibition but not regulated by HyIL-6 treatment (e.g., AQP3 and CCR5); and genes exhibiting a synergic regulation by HyIL-6/CDK8 inhibition combined treatment (e.g., IL-MAP5 and PDCD1) (Figure 6C). Next, we performed, gene set enrichment analysis for evaluating STAT3-mediated transcriptional activity in the absence and presence of CDK8 inhibitor. As expected, genes upregulated by HyIL-6 stimulation were highly enriched in genes known to be upregulated by STAT3 (GEO: GSE21670) (Figure 6D, top panel). Importantly, however, genes upregulated by CDK8 inhibition alone (Figure 6D, middle panel) or in combination with HyIL-6 (Figure 7D, bottom panel) were also highly enriched in known STAT3 targets, indicating that HyIL-6/STAT3 response is mediated by an intrinsic CDK8-dependent axis.

We next asked whether CDK8 could regulate STAT3 binding profiles to chromatin in a genome-wide manner. To assess that question, we carried out STAT3 ChIP-seq in unstimulated or stimulated Th-1 cells with the three conditions described above for 1 h. As expected, in unstimulated Th-1 cells, we detected very low STAT3 DNA binding, which was significantly enhanced upon HyIL-6 treatment (Figure 6E; Figure S7C; Table S3). In Th-1 cells treated with the CDK8 inhibitor alone, we observed levels of STAT3 binding that resembled those obtained in native unstimulated cells (Figure 6E). Strikingly, we observed a synergistic increase in binding intensity of STAT3 across target sites in cells stimulated with the combined HyIL-6/CDK8 inhibitor treatment when compared to HyIL-6 treatment alone, suggesting that inhibition of CDK8 activity amplifies the intensity of HyIL-6-induced STAT3 binding to its target sites (Figure 6E). This increase in STAT3 binding was observed in genes from all three categories described in Figure 6C (Figure 6F). Interestingly, the category of genes not regulated by CDK8 inhibition observed in our RNA-seq study was composed of a set of immediate early genes, such as BCL3 and SOCS3 that are rapidly induced after IL-6 treatment, when compared to HyIL-6 treatment alone, suggesting that inhibition of CDK8 activity led to an average of a 3-fold increase in the number of IL-17-positive cells after 10 days of expansion (Figures 7C and 7D). As expected, we did not see changes in IFNγ expression by the T cells stimulated in the presence of the CDK8 inhibitor (Figures 7B, 7C, and 7E), suggesting a specific regulation of STAT3-mediated Th-17 differentiation by the CDK8 inhibitor. Moreover, CDK8 inhibition resulted in an increase in the levels of secreted IL-17 as measured by ELISA (Figure 7F), but not of IFNγ (Figure 7G). Overall, our data agree with a model where CDK8 modulates STAT3 transcriptional processivity by fine-tuning its gene loci binding dwell-time, leading to a negative regulation of IL-6-mediated activities.

**DISCUSSION**

Our study explores how Ser phosphorylation regulates STAT3 activities in human primary T cells. Overall, our study provides molecular evidence that establishes CDK8 as a master regulator of STAT3 transcriptional activities and presents a potential

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Figure 5. Regulation of the Phosphoproteomic Landscape of IL-6 in Human Primary CD4+ Th-1 Cells by CDK8
(A) Volcano plot of the CDK8-dependent HyIL-6-upregulated phosphosit in human primary Th-1 cells (top panel) and the 24 more affected phosphosit (bottom panel).
(B) Volcano plot of the CDK8-dependent HyIL-6-downregulated phosphosit in human primary Th-1 cells (top panel) and the 24 more affected phosphosit (lower panel). Phosphoproteptides identified in three biological replicates are shown as log-transformed SILAC ratios plotted against log-transformed p values (two-sided t test). Select phosphoproteptides are labeled (see Table S2 for full list). For (A) and (B), phosphosit regulated by HyIL-6 in a CDK8-dependent way and changed more than 1.5-fold with a p value of <0.05 are shown in red (decreased) or blue (increased), and highlighted in dark-gray is the effect of MSC2530818 on those same phosphosit.
(C) GO analysis showing the cellular location of the phosphosit regulated by HyIL-6 in a CDK8-dependent manner.
(D) GO analysis showing the main pathways and cellular processes regulated by HyIL-6 in human primary Th-1 cells.
(E) GO analysis showing the main pathways and cellular processes regulated by HyIL-6 in human primary Th-1 cells in a CDK8-dependent manner.
(F) Pie charts showing the number of HyIL-6-regulated and CDK8-dependent phosphosit involved in the regulation of transcription (left graph) or RNA-Pol-II-mediated transcription (right graph).
(G) The scheme shows the cellular location and molecular function of the proteins regulated by phosphorylation in response to HyIL-6 stimulation in human primary CD4+ Th-1 cells in a CDK8-dependent fashion as determined by DAVID analysis.
A

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B

C

- **BCL3**
  - Unstimulated: 150 ± 10
  - HylL6: 250 ± 20
  - MSC: 200 ± 15
  - HylL6+MSC: 300 ± 25

- **AQP3**
  - Unstimulated: 125 ± 10
  - HylL6: 250 ± 20
  - MSC: 200 ± 15
  - HylL6+MSC: 300 ± 25

- **SOCS3**
  - Unstimulated: 10 ± 1
  - HylL6: 20 ± 2
  - MSC: 15 ± 1.5
  - HylL6+MSC: 30 ± 3

- **CCR5**
  - Unstimulated: 15 ± 1.5
  - HylL6: 25 ± 2
  - MSC: 20 ± 1.5
  - HylL6+MSC: 35 ± 3

- **PDCD1**
  - Unstimulated: 10 ± 1
  - HylL6: 20 ± 2
  - MSC: 15 ± 1.5
  - HylL6+MSC: 30 ± 3

D

- **STAT3 up-regulated genes**
  - Unstimulated
  - HylL6
  - MSC
  - HylL6+MSC

  - **NES**: 1.906
  - **FDR**: 0.0

E

- **STAT3 Binding Intensity**
  - Unstimulated
  - HylL6
  - MSC
  - HylL6+MSC

  - **NES**: 1.849
  - **FDR**: 0.0

F

- **STAT3 Binding genes**
  - Unstimulated
  - HylL6
  - MSC
  - HylL6+MSC

  - **NES**: 1.770
  - **FDR**: 0.0

G

- **STAT3 Binding genes**
  - Unstimulated
  - HylL6
  - MSC
  - HylL6+MSC

  - **NES**: 2.054
  - **FDR**: 0.0

(legend on next page)
strategy to harness the therapeutic potential of cytokines by fine-tuning CDK8 expression levels and activities in different immune cells.

Although a large body of work in the existing literature supports a model in which Ser phosphorylation positively contributes to STAT3 transcriptional activities, our study suggests a negative role of CDK8 activity in STAT3 DNA binding and transcriptional activities in human T cells, mediated at least in part by Ser phosphorylation of STAT3. How can these apparent contradictory observations be reconciled? Previous studies used STAT3 Ser-to-Ala mutants to investigate how Ser phosphorylation regulates STAT3 transcriptional activities (Wakahara et al., 2012; Wen and Darnell, 1997; Wen et al., 1995; Yang et al., 2020). In this context, CDK8 activity remained intact, and therefore, its contribution to STAT3 transcriptional activity was not explored. In agreement with our observations, these studies reported that Ser phosphorylation of STAT3 regulated its chromatin binding dynamics (Wakahara et al., 2012; Yang et al., 2020). STAT3 Ser phosphorylation contributed to destabilization of STAT3 homodimers, resulting in their release from DNA and in STAT3 Tyr dephosphorylation (Yang et al., 2020). However, despite the stronger DNA binding exhibited by the STAT3 S727A mutant, its transcriptional activity was decreased (Wakahara et al., 2012; Wen et al., 1995; Yang et al., 2020). Our study shows that in T cells, CDK8 inhibition prevents STAT3 Ser phosphorylation by IL-6 and increases STAT3 chromatin binding. However, contrary to previous observations, in this context, prolonged STAT3 binding to DNA results in an increased STAT3-dependent transcription. One explanation for these discrepancies in STAT3 activities upon blockage of its Ser phosphorylation could be found in the use of different cell types in the different studies. Although previous studies used cancer cell lines (Wakahara et al., 2012; Wen et al., 1995; Yang et al., 2020), we have used primary human T cells, which thus does not rule out possible differences in the epigenetic landscape between these different cell types that dictates STAT binding profiles. An alternative explanation could be that mutation of S727 in STAT3 and blocking CDK8 activity produce different effects on STAT3 transcriptional activity. Importantly, we show that the STAT3/CDK complex is stabilized in the context of the STAT3 S727A mutant upon IL-6 stimulation. Thus, it is possible that in this context, sustained CDK8 activity, as a result of a stable STAT3/CDK8/DNA complex, could result in transcriptional repression as the one observed for the STAT3 S727A mutant. Our phosphoproteomic study agrees with this model and shows that IL-6 induces the phosphorylation of different transcription factors, including CREM, RelB, and FLI1, in a CDK8-dependent manner, highlighting that active CDK8 elicits a broader regulation of the transcription machinery in response to IL-6 stimulation. Further molecular studies will be required to fully understand how the CDK8/STAT3 interaction ultimately fine-tunes transcriptional output.

IL-6 triggers a strong, albeit transient, phosphorylation of STAT1, but the role that STAT1 phosphorylation plays in fine-tuning IL-6 responses is not clear. STAT1 can form heterodimers with STAT3, having the potential to alter its transcriptional activity (Hirahara et al., 2015). Interestingly, CDK8 phosphorylates both STAT1 and STAT3 (Bancerek et al., 2013), but whether this phosphorylation regulates the formation of STAT1/STAT3 heterodimers is not known at the moment. In addition, STAT1 levels are modulated during inflammation as a consequence of high IFN-α/γ expression in this inflammatory environment, leading to altered responses by cytokines triggering classical STAT3 responses, through the modulation of STAT1/3 homo- and hetero-dimers (Ho and Ivashkiv, 2006). Further studies will be required to address whether CDK8 regulates formation of STAT3 homo- and hetero-dimers and if this regulation is altered in inflammatory environments in which STAT levels are different.

The role that CDK8 plays in transcription regulation has been controversial. Although initially CDK8 was identified as a negative regulator of RNA-Pol II transcription (Jeronimo and Robert, 2017; Knuesel et al., 2009), more recent studies have described showing that CDK8 can also act as a positive modulator of RNA Pol II transcriptional activities (Chen et al., 2017; Donner et al., 2007, 2010). Our results suggest a negative role of CDK8 in STAT3-mediated transcription. CDK8 activity triggers STAT3 dissociation from chromatin and terminates STAT3-mediated transcription. Supporting our model, a series of recent studies have described that modulation of CDK8 activity, by either small-molecule inhibitors or genetic deletion, fine-tune responses elicited by different immune cells. Specific deletion of CDK8 in NK cells results in an enhancement of their anti-tumor responses (Witalisz-Siepracka et al., 2018), an activity that heavily relies on the action of different cytokines on natural killer (NK) cells (Hu et al., 2019). Moreover, small-molecule inhibitors targeting CDK8 promote the differentiation of T reg cells and Th-17 cells when T cells were placed under T reg polarizing conditions (Akamatsu et al., 2019; Guo et al., 2019) or Th-17 polarizing conditions (Figure 7), respectively.
In both instances, the polarizing conditions were enriched in different cytokines, highlighting a potentially broad regulation of cytokine responses by CDK8. Interestingly, CDK8 expres-
sion itself is dynamic. Naive T cells express non-detectable levels of CDK8, which are upregulated upon T cell activation and differentiation (Howden et al., 2019). Thus, it is possible that cells can regulate their CDK8 levels to establish different thresholds of cytokine sensitivity. In agreement with this model, despite high levels of STAT3 Tyr phosphorylation and STAT3 chromatin binding induced by IL-6 treatment, we detected very few genes induced by IL-6 in Th-1 cells, which express high levels of CDK8 (Howden et al., 2019).

**Limitations of Study**

Our study shows that CDK8 regulates STAT3 transcriptional activities by limiting its binding to the target loci. However, whether this repressor effect results from CDK8-mediated phosphorylation of STAT3 on its S727 residue remains an open question. Previous studies described that STAT3 Ser phosphorylation regulates its chromatin binding dynamics, consistent with our model (Wakahara et al., 2012; Yang et al., 2020). However, comparison of chromatin binding profiles between STAT3 WT and its S727A counterpart would be ideal to further support the relative importance of STAT3 Ser phosphorylation in the regulation of STAT3-DNA binding kinetics and its transcriptional activities. Due to the technical challenges in genetic manipulation of primary human T cells, at present, the genomic tools available to examine the role of STAT3 Ser phosphorylation in regulating its transcriptional activities in human primary T cells are inadequate. Furthermore, elucidating whether CDK8 activity regulates the formation of STAT1/STAT3 homo- and hetero-dimers upon IL-6 stimulation,
ultimately impacting IL-6-mediated gene expression programs, remains important.

**STAR★METHODS**

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**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.celrep.2020.108545.

**ACKNOWLEDGMENTS**

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**AUTHOR CONTRIBUTIONS**

J.M.-F. and I.M. conceived the project; J.M.-F., L.W., E.P., A.C., S.W., M.K., S.M., and I.M. wrote the manuscript; J.M.-F., E.P., and S.W. performed signaling and cellular experiments; J.M.-F., L.W., A.C., M.K., S.M., and I.M. designed, performed, and analyzed RNA-seq and ChiP-seq studies.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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**REFERENCES**


CDK8 is a stimulus-specific positive coregulator of p53 target genes. Mol. Cell 27, 121–133.


**STAR METHODS**

**KEY RESOURCES TABLE**

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<td>TOMTOM</td>
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RESOURCE AVAILABILITY

Lead Contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact Ignacio Moraga Gonzalez (imoragagonzalez@dundee.ac.uk).

Materials Availability
This study did not generate new unique reagents.

Data and Code Availability
The phosphoproteomic data have been deposited in the ProteomeXchange: PXD020964 (www.proteomexchange.com). The raw and processed ChIP-seq and RNA-seq data are deposited to GEO: GSE147399.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Human Primary T cells
Peripheral blood mononuclear cells (PBMCs) from healthy donors were purified from buffy coats acquired from the Scottish Blood Transfusion Service and used for signaling and phosphoproteomics experiments.

In vitro polarized human Th-1 cells generated from human PBMCs acquired from StemCell Technologies (Cat#70025) were used for RNA-seq and ChiP-seq experiments.

Cell Lines
Hut78 (cat#TIB-161) and HEK293T (cat#CRL-11268) cells were obtained from ATCC (www.lgcstandards-atcc.org).

METHOD DETAILS

Protein expression and purification
HyIL-6 (Fischer et al., 1997) cloned into the pAcGP67-A vector (BD Biosciences) in frame with an N-terminal gp67 signal sequence and a C-terminal hexahistidine tag was produced using the baculoviral expression system, as previously described (LaPorte et al., 2008). The baculoviral stocks were prepared in Spodoptera frugiperda (Sf9) cells grown in SF900III media (Invitrogen, #12658027) and used to infect Trichoplusiani ni (High Five) cells grown in InsectXpress media (Lonza, #BELN12-730Q) for protein expression. After 48h infection, secreted protein was captured from High Five supernatants using HisPur Ni-NTA resin (Thermo Scientific, #88223) affinity chromatography, concentrated, and purified by size exclusion chromatography on a Enrich SEC 650 1 × 300 column (Biorad), equilibrated in 10 mM HEPES (pH 7.2) containing 150 mM NaCl. Recombinant HyIL-6 was purified to greater than 98% homogeneity.

CD4+ and CD8+ T cell isolation
Peripheral blood mononuclear cells (PBMCs) from healthy donors were purified from buffy coats (Scottish Blood Transfusion Service) by density gradient centrifugation following manufacturer’s instructions (Lymphoprep, StemCell Technologies, #07801). For CD4+ and CD8+ T cells isolation, 1 × 10^8 PBMCs per donor were stained with anti-CD4 FiTC (Biolegend, #357406) or anti-CD8 FITC (Biolegend, #344704) antibodies and isolated by magnetic activated cell sorting (MACS, Miltenyi) using anti-FiTC microbeads (Miltenyi, #130-048-701) according to manufacturer’s instructions to a purity ≥ 99%.

Dose-response and kinetic experiments
For dose-response experiments of STAT1 or STAT3 phosphorylation, 96-well plates were prepared with 30 μL of cells at 5 × 10^5 cells/mL. Cell were then stimulated with different concentrations to obtain the dose-response curves. After stimulation cells were fixed with 2% formaldehyde for 10 minutes at RT.

For kinetics experiments, cell suspensions were stimulated with saturating concentrations of the cytokines (10 nM HyIL-6) as indicated and cells finally fixed with 2% formaldehyde for 10 minutes at RT.

Permeabilization, fluorescence barcoding and antibody staining
After fixation, cells were collected by centrifugation at 1200 rpm for 5 min, formaldehyde blocked by washing the cells with 200 μL of PBS containing 5 mg/ml BSA (PBSA) and collected again by centrifugation at 1200 rpm for 5 min. Then, cells were resuspended and permeabilized in ice-cold methanol for 20 minutes on ice. Cells were then fluorescently barcoded (Krutzik and Nolan, 2006) using a combination of different concentrations of amino-acid reactive dyes (PacificBlue #10163, DyLight800 #46421, Thermo Scientific). Finally, cells were pooled and stained with anti-CD3BV510 (Biolegend, #300448), anti-CD4PE (Biolegend, #357404), anti-CD8AF700 (Biolegend, #300920), anti-pSTAT1-Tyr701AF647 (Cell Signaling, #8009S), anti-pSTAT1-Ser727AF488 (Biolegend, #686410), anti-pSTAT3-Tyr705AF488 (Biolegend, #651006) and anti-pSTAT3-Ser727AF647 (Biolegend, #698914). Cells were analyzed in a CytoFlex.
S flow cytometer (Beckman Coulter) with the individual cell populations being identified by their barcoding pattern and mean fluorescence intensity (MFI) for the different forms of STAT1 or STAT3 measured.

**Phospho-FLOW**

Resting PBMCs isolated as described before fromuffy coats or upon activation forthree days with ImmunoCult Human CD3/CD28 T Cell Activator (StemCell, #10971) following manufacturer instructions in the presence of 20 ng/mL IL2 (Novartis, #709421) were starved for 24 hours in RPMI 1640 (Invitrogen) containing 10% fetal bovine serum (FBS, Invitrogen, #A3160801) and then stimulated with 10nM HyIL-6 or 0.1 ng/mL IL2 (Novartis #709421). Then, cells were fixed with 2% formaldehyde, permeabilized with ice-cold methanol and barcoded as described above. Finally, cells were pooled and stained with anti-CD3ε (Biolegend, #304448), anti-CD4ε (Biolegend, #357404), anti-CD8α (Biolegend, #300920), anti-pSTAT1-Y701(AF647) (Cell Signaling, #8009S), anti-pSTAT1-S727(AF648) (Biolegend, #686410), anti-pSTAT3-Y705(AF648) (Biolegend, #651006) and anti-pSTAT3-S727(AF647) (Biolegend, #698913), anti-pSTAT4-Y693(AF648) (BD Biosciences, #558136), anti-pSTAT5-Y694(AF647) (Cell Signaling, #9365S), anti-pSTAT6-Y641(AF488) (BD Biosciences, #612600), anti-pERK-T202/Y204(AF488) (eBiosciences, #560392) and anti-PLCγ(AF488) (BD Biosciences, #8677S), anti-IRF1(AF647) (Cell Signaling, #14105S), anti-IRF4(AF647) (Biolegend, #646408), anti-IRF7(AF647) (Biolegend, #650007), anti-GATA3(AF488) (Biolegend, #653807), anti-STAT4(AF488) (Biolegend, #359707), anti-MYC(AF488) (Cell Signaling, #12855S), anti-O-GlcNAc(AF488) (NOVUS Biologicals, #NB300-524AF647), anti-STAT5b(AF647) (Cell Signaling, #9716S), anti-pAKT-S473(AF488) (Cell Signaling, #4071S), anti-pAKT-T308(AF647) (Cell Signaling, #82975S), anti-pCREB-S133(AF488) (Cell Signaling, #9187S), anti-pS6K1(S240/244)(AF647) (Cell Signaling, #14323S), anti-pERK-T202/Y204(AF488) (eBiosciences, #53-9109-41), anti-pAKT-S473(AF488) (Cell Signaling, #4071S), anti-p-AKT-T308(AF647) (Cell Signaling, #82975S), anti-pCREB-S133(AF488) (Cell Signaling, #9187S), anti-pS6K1(S240/244)(AF647) (Cell Signaling, #14323S), anti-pERK-T202/Y204(AF488) (eBiosciences, #53-9109-41), anti-pAKT-S473(AF488) (Cell Signaling, #4071S), anti-p-AKT-T308(AF647) (Cell Signaling, #82975S), anti-pCREB-S133(AF488) (Cell Signaling, #9187S), anti-pS6K1(S240/244)(AF647) (Cell Signaling, #14323S), anti-pERK-T202/Y204(AF488) (eBiosciences, #53-9109-41), anti-pAKT-S473(AF488) (Cell Signaling, #4071S), anti-p-AKT-T308(AF647) (Cell Signaling, #82975S), anti-pCREB-S133(AF488) (Cell Signaling, #9187S), anti-pS6K1(S240/244)(AF647) (Cell Signaling, #14323S), anti-pERK-T202/Y204(AF488) (eBiosciences, #53-9109-41), anti-pAKT-S473(AF488) (Cell Signaling, #4071S), anti-p-AKT-T308(AF647) (Cell Signaling, #82975S), anti-pCREB-S133(AF488) (Cell Signaling, #9187S), anti-pS6K1(S240/244)(AF647) (Cell Signaling, #14323S), anti-pERK-T202/Y204(AF488) (eBiosciences, #53-9109-41), anti-pAKT-S473(AF488) (Cell Signaling, #4071S), anti-p-AKT-T308(AF647) (Cell Signaling, #82975S), anti-pCREB-S133(AF488) (Cell Signaling, #9187S), anti-pS6K1(S240/244)(AF647) (Cell Signaling, #14323S), anti-pERK-T202/Y204(AF488) (eBiosciences, #53-9109-41), anti-pAKT-S473(AF488) (Cell Signaling, #4071S), anti-p-AKT-T308(AF647) (Cell Signaling, #82975S), anti-pCREB-S133(AF488) (Cell Signaling, #9187S), anti-pS6K1(S240/244)(AF647) (Cell Signaling, #14323S), anti-pERK-T202/Y204(AF488) (eBiosciences, #53-9109-41).
Phosphopeptide enrichment in the peptide fractions generated as described above was carried out using MagResyn Ti-IMAC following manufacturer instructions (2BScientific, MR-TIM002). Phosphopeptide samples were analyzed using a nanoflow liquid chromatography system (Ultimate 3000 RSLCnano system, Thermo Scientific) coupled to a Q Exactive Plus Mass Spectrometer (Thermo Scientific). Samples (10 μl) were loaded onto a C18 trap column and washed for 5 minutes with 0.1% formic acid. Peptides were resolved using a gradient (170 min, 0.3 μl/min) of buffer A (0.1% formic acid) and buffer B (80% acetonitrile in 0.08% formic acid): 5% buffer B for 5 min, 5%–35% buffer B for 125 min, 35%–98% buffer B for 2 min, 98% buffer B for 20 min, 98%–2% buffer B for 1 min and 2% buffer B for 17 min. Peptides, initially trapped on an Acclaim PepMap 100 C18 column (100 μm x 2 cm, Thermo Scientific), were separated on an Easy-Spray PepMap RSLC C18 column (75 μm x 50 cm, Thermo Scientific), and finally transferred to a Q Exactive Plus Mass Spectrometer via an Easy-Spray source with temperature set at 50 °C and a source voltage of 2.0 kV. For the identification of peptides, a top 15 method (1 MS plus 15 MS², 150 min acquisition) consisting of full scans and mass range (m/z) between 350 to 1600 (m/z) for MS search and 200 to 2000 (m/z) for MS² search was used. For the MS scan the Q Exactive Plus Mass Spectrometer was operated in a data dependent acquisition mode, resolution of 70,000 with a lock mass set at 445.120024 and max fill time of 20 ms. For the MS² scan Q Exactive Plus Mass Spectrometer was operated in a centroid mode, resolution of 15,000, with isolation window = 1.4 (m/z), normalized collision energy = 27, max fill time of 100 ms and dynamic exclusion of 45.0 s.

Inhibition of Ser727 STAT3 phosphorylation

Resting CD4⁺ T cells were labeled with anti-CD4-FITC antibody (Biolegend, #357406) and isolated from human PBMCs by magnetic activated cell sorting (MACS, Miltenyi) using anti- FITC microbeads (Miltenyi, Cat#130-048-701) following manufacturer instructions. Then, CD4⁺ T cells were activated for three days with ImmunoCult Human CD3/CD28 T Cell Activator (StemCell, #10971) following manufacturer instructions in the presence of 20 ng/mL IL2 (Novartis, #709421). After activation, cells were expanded for 5 days in the presence of IL-2, these cells were then either not-stimulated (control) or stimulated with IL-6 or different IL-6 variants for 1 hr, cells were then immediately fixed with 1% methanol-free formaldehyde (Formaldehyde 16%, Methanol-Free, Fisher Scientific, PA, USA) and allowed to lyze in ice cold PBS then analyzed by flow cytometry. Alternatively, these cells were polarized human Th-1 cells generated from human PBMCs (StemCell Technologies, Cat#70025) were expanded in the presence of 20 ng/mL IL2 for 10 days and cells were then washed with complete media and rested for 24 hr starvation in the absence of IL-2, these cells were then either not-stimulated (control) or stimulated with IL-6 or different IL-6 variants for 1 hr, cells were then immediately fixed with 1% methanol-free formaldehyde (Formaldehyde 16%, Methanol-Free, Fisher Scientific, PA, USA) and allowed to lyze in ice cold PBS then analyzed by flow cytometry.

Chromatin immunoprecipitation by sequencing (ChIP-Seq)

In vitro polarized human Th-1 cells generated from human PBMCs (StemCell Technologies, Cat#70025) were expanded in the presence of IL-2 for 10 days and cells were then washed with complete media and rested for 24 hr starvation in the absence of IL-2, these cells were then either not-stimulated (control) or stimulated with IL-6 or different IL-6 variants for 1 hr, cells were then immediately fixed with 1% methanol-free formaldehyde (Formaldehyde 16%, Methanol-Free, Fisher Scientific, PA, USA) and allowed to lyze in ice cold PBS then analyzed by flow cytometry.
at room temperature for 10 min with gentle rocking cells were then washed twice with cold PBS. For each STAT3 ChIP-seq library sample, approximately 10 × 106 cells were used and the fixed cell pallettes were kept at −80°C prior to further processing. The ChIP-seq experiments were performed as previously described (Martinez-Fabregas et al., 2019) with some modification as described below. In brief, the frozen cell pellets were thawed on ice and washed once with 1 mL cold PBS by centrifugation at 5000 RPM for 5 min, the resulting cell pellets were resuspended in 500 uL of lysis buffer (1X PBS, 0.5% Triton X-100, complete EDTA-free protease inhibitor cocktail, Roche Diagnostics, Basel, Switzerland) and incubated for 10 min on ice, followed by a 5 min centrifugation at 5000 RPM. Then the pellets were washed once with 1 mL of sonication buffer (1X TE, 1: 100 protease inhibitor cocktail), re-suspended in 750 uL of sonication buffer (1X TE, 1: 100 protease inhibitor cocktail and 0.5 mM PMSF) and sonicated for 20 cycles (on-20 sec and off-45 sec) on ice using VCX-750 Vibra Cell Ultra Sonic Processor (Sonic, USA). The sonicated lysates were centrifuged 20 min at 14000 RPM and the clear lysate supernatants were collected and incubated with 30 uL of Protein-A Dynabeads (ThermoFisher, USA) that were pre-incubated with 10 uM of anti-STAT3 antibody (anti-Stat3, 12640S, Cell Signaling Technology) at 4°C overnight with gentle rotation. Next day, the beads were washed 2 times with RIPA-140 buffer (0.1% SDS, 1% Triton X-100, 1 mM EDTA, 10 mM Tris pH 8.0, 300 mM NaCl, 0.1% NaDOC), 2 times with RIPA-300 buffer (0.1% SDS, 1% Triton X-100, 1 mM EDTA, 10 mM Tris, 300 mM NaCl, 0.1% NaDOC), 2 times with LiCl buffer (0.25 mM LiCl, 0.5% NP-40, 1 mM EDTA, 10 mM Tris pH 8.0, 0.5% NaDOC), once with TE-0.2% Triton X-100 and once with TE buffer. Crosslinks were reversed by incubating the bound complexes in 60 uL containing 4.5 uL of 10% SDS and 7.5 uL of 20 mg/mL of proteinase K (Thermofisher, USA) at 65°C overnight for input samples, we used 6 uL of 10% SDS and 10 uL of 20 mg/mL of proteinase K. Then, the supernatants were collected after a magnet and beads were further washed using one in TE 0.5M NaCl buffer. Both supernatants were combined, and DNA was extracted with phenol/chloroform, followed by precipitation with ethanol and re-suspended in TE buffer. The library was constructed following the manufacturer protocol of the KAPA LTP Library Preparation Kit (KAPA Biosystems, Roche, Switzerland). ChIP DNA libraries were ligated with the Bioo scientific barcoded adaptors (BIOO Scientific, Perkin Elmer, USA) with T4 DNA ligase according to KAPA LTP library preparation protocol and the ligated ChIP DNA libraries were purified with 1.8x vol. Agencourt AMPure XP beads and PCR amplified using KAPA hot start High-Fidelity 2X PCR Master Mix and NextFlex index primers (Bioo Scientific, PerkinElmer) for 12 cycle by following thermocycler cycles: 30 s hot start at 98°C, followed by 12 cycle amplification [98°C for 10 s, 60°C for 30 s and 72°C for 30 s] and final extension at 72°C for 1 min. The amplification and quality of the ChIPseq libraries were checked by running 10% of the samples in E-Gel Agarose Gels with SYBR Safe DNA Gel Stain (ThermoFisher Scientific, USA), and if necessary, samples were re-amplified additional four cycles using the same thermocycler protocol described above. Then, the libraries were purified and size-selected using Agencourt AMPure XP beads (1.25x vol. to remove short fragments. The concentration of ChIP-DNA libraries was measured by Qubit-4 fluorometer (ThermoFisher, USA) and equal amounts of each sample were pooled and 50 bp paired-end reads were sequenced on an Illumina 4000 platform by GENEWIZ technology (GENEWIZ, USA).

RNA-sequencing
For RNA-seq library preparation, in vitro polarized human Th-1 cells generated from human PBMCs (StemCell Technologies, Cat#70025) either not stimulated or stimulated with HyIL-6 in the presence or absence of 2 μM MSC2530818 variants at 37°C for 6 hr, total RNA was extracted and RNaseq libraries were prepared by Edinburg Sequencing Core facility.

CDK8 and CDK9 knock-down in HEK293T cells
CDK8 and CDK9 knock-down in HEK293T cells was done using Lipofectamine RNAiMAX following manufacturer instructions. Briefly, cells were seeded at 6x10^5 per well in 6-well plates and transfected with siRNA SMARTPools against CDK8 (Dharmacon, Cat# LQ-003242-00-0005) and/or CDK9 (Dharmacon, Cat# LQ-003242-00-0005) 24 hours later. Cells were cultured for 48 hours in the presence of the siRNA mixture before using them for FACS or western blot analysis. Knock-down efficiency was checked by western blotting using anti-CDK8 (G398, Cat# 4101), anti-CDK9 (C12F7, Cat# 2316) and anti-GAPDH (14C10, Cat# 2118).

In vitro CDK assay
In vitro CDKs phosphorylation reactions (20 μL total volume) were as follows: 10 μL of 2x CDK phospho assay buffer (50mM β-glycerophosphate pH 7.4, 10mM MgCl2, 10mM NaF, 1mM DTT) with 100 μM ATP and 5 μL of STAT3 at 20ng/μL. Reactions were initiated by adding 5 μL of varying amounts of CK7, CK8 or CK9. Reactions were incubated for 30 minutes at 30°C and stopped by adding 5 μL of 4x SDS sample buffer and heating to 95°C for 5 minutes. All kinase reactions were performed at least three times. ATP was purchased from Sigma (Cat# A2383-10G), human recombinant CDKs were purchased from Thermo (CDK7/CyclinH/MNAT1 Cat# PV3868, CDK8/CyclinC Cat# PV4402 and CDK9/CyclinK Cat# PV4335) and human recombinant STAT3 was purchased from Novus-Biologicals (Cat# H0000677-P01-10 μg).

Crispr/Cas9 generation of STAT3 KnD Hut78 cells
5 μl of 200 μM Alt-R CRISPR-Cas9 crRNA (IDT, Hs.Cas9.STAT3.1.AF) were added to 5 μL of 200 μM Alt-R CRISPR-Cas9 tracrRNA (IDT, #1072532), heated to 95°C for 5 min and finally the tube cooled to room temperature. Then, 1.2 μL RNA duplex was mixed with 1.7 μL Alt-R S.p. HiFi Cas9 Nuclease V3 (IDT, #1081059) and 2.1 μL sterile PBS and incubated at RT for 20 min, 2 x 105 Hut78 cells resuspended in 8 μL of buffer R (Next Generation Transfection System, Thermo) were added to the tubes with the RNP complexes. The
electroporation parameters used were: three pulses of 1,325 V with a pulse width of 10 ms. Then, reactions were added directly into antibiotic-free media in a well of the 96-well plate and incubated at 37°C for 16 hr. HeLa cells electroporated with RNP particles were transferred into IMDM media containing 10% FCS and Pen/Strep, expanded and finally individual clones were isolated and tested for STAT3 expression levels.

**STAT3 KnD Hut78 Cells**

**STAT3 Reconstitution**

STAT3 KnD Hut78 cells were reconstituted with a plasmid coding for STAT3 WT or STAT3 S727A mutants by electroporation. 2 x 10^7 cells were resuspended in 0.25 mL of Ingenio electroporation solution (Mirus, #MIR50111), 30 µg of the appropriate construct added and the mix transferred into 4 mm gap-width cuvettes and incubated at RT for 15 minutes. Cells were electroporated using the Bio-Rad X-Gene Pulser system (0.28 kV, 960 µF), cells transferred to pre-warmed media without Pen/Strep and allow to recover for 24 hours.

**T cells population differentiation**

Resting CD4+ T cells isolated as described above were activated under Th-1 or Th-17 polarizing conditions. Briefly, resting human CD4+ T cells freshly isolated were activated using ImmunoCult Human CD3/CD28 T Cell Activator (StemCell, Cat#10971) following manufacturer instructions for 3 days in the presence of the cytokines required for the different CD4+ T cells populations: Th-1 (IL-2 (20 ng/ml), anti-IL-4 (10 ng/ml, BD Biosciences, Cat#554481), IL-12 (20 ng/ml, R and D Systems, Cat#201-LB/CF), IL-23 (10 ng/ml, R and D Systems, Cat#1290-IL), anti-IL-4 (10 ng/ml, BD Biosciences, Cat#554481), anti-IFNγ (10 ng/ml, BD Biosciences, Cat#554698)). After three days of priming, cells were expanded in the presence of IL-2 (20 ng/ml), Th-1 and Th-17 cells were restimulated for 6 hr in the presence of PMA (100 ng/ml, Sigma, Cat#P8139), Ionomycin (1 µM, Sigma, I0634) and Brefeldin A (5 µg/ml, Sigma, B7651) before FACS analysis. In all cases cells were fixed with 2% formaldehyde and prepared to be analyzed by FACS. Cells were then permeabilised with Saponin 2% in PBS for 20 min at room temperature and then stained in Saponin 2% in PBS with the appropriate antibodies: Th-1 ((anti-CD3 BV510 (1:100, Biolegend, Cat#300448), anti-CD4 PE (1:100, Biolegend, Cat#357404), anti-CD8 AF700 (1:100, Biolegend, Cat#300920), anti-IFNγ AF488 (1:100, Biolegend, Cat#512334)) and analyzed in a CytoFLEX S (Beckman Coulter).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

**Bioinformatics**

The following software were used:

- MaxQuant (Cox and Mann, 2008)
- Andromeda (Cox et al., 2011)
- DAVID GO analysis tool (Huang et al., 2007, 2009)
- FastQC v0.11.8 (www.bioinformatics.babraham.ac.uk)
- RSEM v1.3.1 (Li and Dewey, 2011)
- edgeR v3.24.0 (Robinson et al., 2010)
- Datagraph v4.5 (www.visualdatatools.com)
- PRISM v8.4.0 (https://www.graphpad.com/scientific-software/prism/)
- GSEA v4.0.3 (Subramanian et al., 2005)
- Metascape (Zhou et al., 2019)
- Bowtie v1.2.2 (Langmead et al., 2009)
- SameTools v1.9 (Li et al., 2009)
- bamCoverage v3.2.0 (Ramirez et al., 2016)
- MEME Suite v5.0.2 (Bailey et al., 2009)
- TOMTOM (Gupta et al., 2007)
- BEDTools (Quinlan and Hall, 2010)
- UCSC bigWigAverageOverBed v2 (https://genome.ucsc.edu)
- HOMER v4.10 (Gupta et al., 2007)

**Mass spectrometry data analysis**

Q Exactive Plus Mass Spectrometer .RAW files were analyzed, and peptides and proteins quantified using MaxQuant (Cox and Mann, 2008), using the built-in search engine Andromeda (Cox et al., 2011). All settings were set as default, except for the minimal peptide length of 5, and Andromeda search engine was configured for the UniProt Homo sapiens protein database (release date: 2018_09). Peptide and protein ratios only quantified in at least two out of the three replicates were considered, and the p values were determined by Students t test and corrected for multiple testing using the Benjamin-Hochberg procedure (Benjamini and Hochberg, 1995).
DAVID GO ANALYSIS TOOL (HUANG ET AL., 2007, 2009) WAS USED TO FIND STATISTICALLY OVER-REPRESENTED GENE ONTOLOGY (GO) CATEGORIES IN THE PROTEOMIC DATA

RNA-Seq analysis
The quality of libraries was inspected by FastQC v0.11.8. The expression level of mRNA in each library was quantified by ‘rsem-calculate-expression’ in RSEM v1.3.1 (Li and Dewey, 2011) using default parameters except ‘–bowtie-n 1–bowtie-m 100–seed-length 28–paired-end’. The bowtie index required by RSEM software was generated by ‘rsem-prepare-reference’ on all RefSeq genes, downloaded from UCSC table browser on April 2017. edgeR v3.24.0 (Robinson et al., 2010) package was used to normalize gene expression among all libraries and identify differentially expressed genes among samples with following constraints: fold change ≥ 1.5, p value ≤ 0.05. Scatter and bar plots were drawn by Datagraph v4.5 and PRISM v8.4.0, respectively. Geneset enrichment analysis was performed by GSEA v4.0.3 (Subramanian et al., 2005) with default parameters except ‘–collapse No_Collapse -permute gene_set’. Pathway analysis of differentially expressed genes was performed by Metascape (Zhou et al., 2019) on all GO terms related to biological processes, KEGG Pathways, BioCarta Gene Sets and Hallmark Gene Sets.

ChIP-Seq analysis
The quality of libraries was inspected using FastQC v0.11.8. All sequencing reads were aligned to human reference genome (GRCh37; hg19) using bowtie v1.2.2 (Langmead et al., 2009) with default pair-end alignment settings and additional parameters ‘–chunkmbs 1000 S -m 1’. The index for reference genome was constructed by using ‘bowtie-build’ with default parameters. Sorting and indexing of the aligned reads were conducted by Samtools v1.9 (Li et al., 2009). The genome-wide binding profiles (i.e., bigWig files) were generated by bamCoverage v3.2.0 (Ramírez et al., 2016) using parameters ‘–normalizeUsing BPM–minMappingQuality 30–ignoreDuplicates–extendReads 250–blackListFileName hg19.blacklist.bed’. The binding profiles were visualized using IGV genome browser v2.7.0 (Robinson et al., 2011). Binding peaks were called by ‘callpeaks’ procedure from MACS2 v2.1.2 (Zhang et al., 2008) using default parameters except ‘-f BAMPE-nomodel -t treatment -c input’. The identified peaks were further screened against ‘hg19 blacklisted’ genomic regions, mitochondrial DNA, and pseudo-chromosomes. De novo motif findings were performed in 200 bp surrounding the summit of n = 500 top bound regions using MEME Suite v5.0.2 (Bailey et al., 2009) with default parameters except ‘–maxsize 1000000 -dna -mod zoops -nmotifs 10’. De novo motifs were compared against all JASPAR known motifs by TOMTOM (Gupta et al., 2007). STAT3 shared bound regions in HyIL6 (n = 540) or HyIL-6+MSC (n = 2585) stimulated cells were generated by the intersection between bound regions from n = 3 donor using BEDTools (Quinlan and Hall, 2010). STAT3 binding intensity in shared bound regions was calculated by UCSC bigWigAverageOverBed v2 with default parameters and the mean signal intensity was visualized by PRISM v8.4.0. The shared STAT3 bound regions were annotated with the nearest gene by ‘annotatePeaks’ from HOMER v4.10 (Gupta et al., 2007), yielding 475 unique genes. Statistical analyses were performed using the Two-tailed parametric and non-parametric tests as appropriate.

Statistical analysis
Statistical significance of differential induction of phosphopeptide ratios only quantified in at least two out of the three replicates were considered, and the p values were determined by Students t test and corrected for multiple testing using the Benjamini–Hochberg procedure (Benjamini and Hochberg, 1995). For RNA-seq, statistical significance was calculated by edgeR v3.24.0 (Robinson et al., 2010) package. Calculation of statistical significance for binding intensity of STAT3 ChIP-seq was conducted by non-parametric two-tailed Wilcoxon test. Further information related to the statistical analyses performed is provided in the figure legends.