1	Non-canonical role of STAT2 and IRF9 in the regulation of a STAT1-
2	independent antiviral and immunoregulatory transcriptional program
3	induced by IFN β and TNF α
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1 Significance

2 IFN β exhibit potent anti-proliferative, antiviral and immunoregulatory functions. The response to 3 IFN β is context-dependent and is prone to crosstalk with other cytokines, including TNF α . Elevated 4 levels of both IFN β and TNF α are notably observed during infections or chronic autoimmune and 5 inflammatory diseases. A synergistic action of IFN β and TNF α has previously been described, but 6 the underlying mechanism remained elusive. Formation of ISGF3 (STAT1/STAT2/IRF9) is 7 considered a hallmark of the type I IFN response, and the requirement of STAT1 in a specific setting 8 has become a marker of the engagement of type I IFN signaling. The finding that in the presence of 9 TNF α STAT1 is not required to drive a delayed antiviral and immunoregulatory transcriptional 10 program challenges this paradigm. Similarly, the observation that STAT2 and IRF9 act 11 independently to regulate a distinct subset of genes unveil the existence of previously unrecognized 12 pathways that mediate the synergistic action of IFN β and TNF α .

13

14 ABSTRACT

15 IFN β typically induces an antiviral and immunoregulatory transcriptional program through the 16 activation of ISGF3 (STAT1, STAT2 and IRF9) transcriptional complexes. The response to IFNB is 17 context-dependent and is prone to crosstalk with other cytokines, such as TNF α . IFN β and TNF α synergize to drive a specific delayed transcriptional program. Previous observation led to the 18 19 hypothesis that an alternative STAT1-independent pathway involving STAT2 and IRF9 might be 20 involved in gene induction by the combination of IFN β and TNF α . Using genome wide 21 transcriptional profiling by RNASeq, we found that the costimulation with IFNB and TNFa induces a 22 broad antiviral and immunoregulatory transcriptional program independently of STAT1. Additionally, STAT2 and IRF9 are involved in the regulation of only a subset of these STAT1-23 24 independent genes. Consistent with the growing literature, STAT2 and IRF9 act in concert to regulate

1 a subgroup of these genes. Unexpectedly, STAT2 and IRF9 were also engaged in specific 2 independent pathways to regulate distinct sets of IFN β and TNF α -induced genes. Altogether these 3 observations highlight the existence of distinct previously unrecognized non-canonical STAT1-4 independent, but STAT2 and/or IRF9-dependent pathways in the establishment of a delayed antiviral 5 and immunoregulatory transcriptional program in conditions where elevated levels of both IFN β and 6 TNF α are present.

1 INTRODUCTION

2 Interferon (IFN) β plays a critical role in the first line of defense against pathogens, 3 particularly viruses, through its ability to induce a broad antiviral transcriptional response in virtually 4 all cell types (1). IFN β also possesses key immunoregulatory functions that determine the outcome of the adaptive immune response against pathogens (1, 2). IFN β acts through binding to the IFNAR 5 6 receptor (IFNAR1 and IFNAR2) leading to Janus kinases (JAK), JAK1 and Tyk2, mediated 7 phosphorylation of signal transducer and activator of transcription (STAT) 1 and STAT2, and to a 8 lesser extent other STAT members in a cell-specific manner (3, 4). Phosphorylated STAT1 and 9 STAT2 together with IFN Regulatory Factor (IRF) 9 form the IFN-stimulated gene factor 3 (ISGF3) 10 complex that binds to the consensus IFN-stimulated response element (ISRE) sequences in the 11 promoter of hundreds of IFN stimulated genes (ISGs) (5). Formation of the ISGF3 complex is 12 considered a hallmark of the engagement of the type I IFN response, and consequently the 13 requirement of STAT1 in a specific setting has become a marker of the engagement of type I IFN 14 signaling (3, 6). However, in recent years this paradigm has started to be challenged with 15 accumulating evidence demonstrating the existence of non-canonical JAK-STAT signaling mediating 16 type I IFN responses (4, 7).

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18 Over the years, *in vitro* and *in vivo* studies aimed at characterizing the mechanisms and the 19 functional outcomes of IFN β signaling were mostly performed in relation to single cytokine 20 stimulation. However, this unlikely reflects physiological settings, as a plethora of cytokines is 21 secreted in a specific situation. As a consequence, a cell rather simultaneously responds to a cocktail 22 of cytokines to foster the appropriate transcriptional program. Response to IFN β is no exception and 23 is very context-dependent, particularly regarding the potential cross-talk with other cytokines. IFN β 24 and TNF α exhibit context-dependent cross-regulation, but elevated levels of both cytokines are

found during the host response to pathogens, including virus and bacteria, and also in autoinflammatory and autoimmune diseases (8). While the cross-regulation of IFN β and TNF α is well studied, the functional cross-talk between these two cytokines remains poorly known and is limited to the description of a synergistic interaction (9-12). Indeed, costimulation with IFN β and TNF α was found to drive a specific delayed transcriptional program composed of genes that are either not responsive to IFN β or TNF α separately or are only responsive to either one of the cytokine

7 (10, 11).

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9 The signaling mechanisms engaged downstream of the costimulation with IFN β and TNF α 10 remained elusive, but it is implicitly assumed that the fate of the gene expression response requires 11 that both IFN β and TNF α induced signaling pathways exhibit significant cross-talk. Analysis of the 12 enrichment of specific transcription factors binding sites in the promoters of a panel of genes 13 synergistically induced by IFN β and TNF α failed to give a clue about the specificity of the 14 transcriptional regulation of these genes (10). Recently, analysis of the induction of DUOX2 and 15 DUOXA2 genes, which belong to the category of delayed genes that are remarkably induced to high 16 levels in response to the combination of IFN β and TNF α , led to the hypothesis that STAT2 and IRF9 17 activities might segregate in an alternative STAT1-independent pathway that could be involved in 18 gene induction downstream of IFN β and TNF α (12). Further validation was awaited to confirm the 19 existence of this STAT1-independent response and the extent to which it is involved in the regulation 20 of a specific delayed transcriptional program induced by the combination of IFN β and TNF α .

21

In the present study, we aimed to characterize the transcriptional profile of the delayed response to IFN β and TNF α in the absence of STAT1 and evaluate the role of STAT2 and IRF9 in the regulation of this responseTaking advantage of STAT1-deficient cells, we found that the synergistic action of IFN β and TNF α induces a broad delayed antiviral and immunoregulatory transcriptional program independently of STAT1. We also report that STAT2 and IRF9 are differentially involved in the regulation of distinct subsets of genes induced by IFN β and TNF α While IFN β and TNF α act in part through the concerted action of STAT2 and IRF9, specific sets of genes were only regulated either by STAT2 or IRF9. These findings highlight the existence of distinct previously unrecognized non-canonical STAT2 and/or IRF9-dependent pathways that mediate the synergistic action of IFN β and TNF α .

8

9 MATERIAL AND METHODS

10

11 Cell culture and stimulation

12 A549 cells (American Type Culture Collection, ATCC) were grown in F-12 nutrient mixture (Ham) 13 medium supplemented with 10% heat-inactivated fetal bovine serum (HI-FBS) and 1% L-glutamine. 14 The 2ftGH fibrosarcoma cell line and the derived STAT1-deficient U3A cell line, a generous gift 15 from Dr. G. Stark, Cleveland, USA (13), were grown in DMEM medium supplemented with 10% 16 HI-FBS or HI-Fetal Clone III (HI-FCl) and 1% L-glutamine. U3A cells stably expressing STAT1 17 were generated by transfection of the STAT1 alpha flag pRc/CMV plasmid (Addgene plasmid 18 #8691; a generous gift from Dr. J. Darnell, Rockfeller University, USA (14, 15)) and selection with 19 800µg/ml Geneticin (G418). Monoclonal populations of U3A stably expressing STAT1 cells were 20 isolated. A pool of two clones, referred to as U3A-STAT1, was used in the experiments to mitigate 21 the clonal effects. U3A-STAT1 cells were maintained in culture in DMEM supplemented with 10% 22 HI-FCl, 1% Glu, and 200µg/ml G418. All cell lines were cultured without antibiotics. All media and 23 supplements were from Gibco, with the expection of HI-FCl, which was from HyClone.

Mycoplasma contamination was excluded by regular analysis using the MycoAlert Mycoplasma
 Detection Kit (Lonza). Cells were stimulated with IFNβ (1000 U/mL, PBL Assay Science), TNFα
 (10 ng/mL, R&D Systems) or IFNβ (1000 U/mL) +TNFα (10 ng/mL) for the indicated times.

Τ.

5 siRNA Transfection

The sequences of non-targeting control (Ctrl) and STAT2- and IRF9-directed RNAi oligonucleotides
(Dharmacon, USA) have previously been described in (12). U3A cells at 30% confluency were
transfected using the Oligofectamine transfection reagent (Life technologies). RNAi transfection was
pursued for 48 h before stimulation.

10

11 Immunoblot analysis

12 Cells were lysed on ice using Nonidet P-40 lysis buffer as fully detailed in (16). Whole-cell extracts 13 (WCE) were quantified using the Bradford protein assay (Bio-Rad), resolved by SDS-PAGE and 14 transferred to nitrocellulose membrane before analysis by immunoblot. Membranes were incubated 15 with the following primary antibodies, anti-actin Cat #MAB1501 from Millipore, anti-IRF9 Cat 16 #610285 from BD Transduction Laboratories, and anti-STAT1-P-Tyr701 Cat #9171, anti-STAT2-P-17 Tyr690 Cat #4441, anti-STAT1 Cat #9172, anti-STAT2 Cat #4594, all from Cell Signaling, before 18 further incubation with horseradish peroxidase (HRP)-conjugated secondary antibodies (KPL or 19 Jackson Immunoresearch Laboratories). Antibodies were diluted in PBS containing 0.5% Tween and 20 either 5% nonfat dry milk or BSA. Immunoreactive bands were visualized by enhanced 21 chemiluminescence (Western Lightning Chemiluminescence Reagent Plus, Perkin-Elmer Life 22 Sciences) using a LAS4000mini CCD camera apparatus (GE healthcare).

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- 24

1 RNA isolation and qRT-PCR analyses

2 Total RNA was prepared using the RNAqueous-96 Isolation Kit (Ambion) following the 3 manufacturer's instructions. Total RNA (1µg) was subjected to reverse transcription using the 4 QuantiTect Reverse Transcription Kit (Qiagen). Quantitative PCR were performed using either Fast 5 start SYBR Green Kit (Roche) for Mx1, IDO, APOBEC3G, CXCL10, NOD2, PKR, IRF1, and IFIT1 6 and IL8 or TaqMan Gene Expression Assays (Applied Biosystems) for DUOX2, IFI27, SERPINB2, 7 IL33, CCL20, ISG20, OAS3, GRB14, CLEC5A, DGX58, ICAM1, IL1B, JUNB, MMP9, SERPINA1 8 and HERC6. Sequences of oligonucleotides and probes used in PCR reactions are described in 9 Supplemental Table S4. Data collection was performed on a Rotor-Gene 3000 Real Time Thermal 10 Cycler (Corbett Research). Gene inductions were normalized over S9 levels, measured using Fast 11 start SYBR Green Kit or TaqMan probe as necessary. Fold induction of genes was determined using 12 the $\Delta\Delta$ Ct method (17). All qRT-PCR data are presented as the mean \pm SEM.

13

14 **RNA-sequencing (RNASeq)**

15 Total RNA prepared as described above was quantified using a NanoDrop Spectrophotometer ND-16 1000 (NanoDrop Technologies, Inc.) and its integrity was assessed using a 2100 Bioanalyzer 17 (Agilent Technologies). Libraries were generated from 250 ng of total RNA using the NEBNext 18 poly(A) magnetic isolation module and the KAPA stranded RNA-Seq library preparation kit (Kapa 19 Biosystems), as per the manufacturer's recommendations. TruSeq adapters and PCR primers were 20 purchased from IDT. Libraries were quantified using the Quant-iTTM PicoGreen® dsDNA Assay Kit 21 (Life Technologies) and the Kapa Illumina GA with Revised Primers-SYBR Fast Universal kit (Kapa 22 Biosystems). Average size fragment was determined using a LabChip GX (PerkinElmer) instrument. 23 Massively parallel sequencing was carried out on an Illumina HiSeq 2500 sequencer. Read counts 24 were obtained using HTSeq. Reads were trimmed from the 3' end to have a Phred score of at least 30.

1 Illumina sequencing adapters were removed from the reads and all reads were required to have a 2 length of at least 32bp. Trimming and clipping was performed using Trimmomatic (18). The filtered 3 reads were aligned to the Homo-sapiens assembly GRCh37 reference genome. Each readset was 4 aligned using STAR (19) and merged using Picard (http://broadinstitute.github.io/picard/). For all 5 samples, the sequencing resulted in more than 29 million clean reads (ranging from 29 to 44 million 6 reads) after removing low quality reads and adaptors. The reads were mapped to the total of 63679 7 gene biotypes including 22810 protein-coding genes. The non-specific filter for 1 count-per million 8 reads (CPM) in at least three samples was applied to the reads and 14,254 genes passed this criterion. 9 The entire set of RNAseq data has been submitted to the Gene Expression Omnibus (GEO) database 10 (http://www.ncbi.nlm.nih.gov/geo) under accession number GSE111195.

11

12 **Bioinformatics analysis**

13 Differential transcripts analysis. A reference-based transcript assembly was performed, which allows 14 the detection of known and novel transcripts isoforms, using Cufflinks (20), merged using 15 Cuffmerge (cufflinks/AllSamples/merged.gtf) and used as a reference to estimate transcript 16 abundance and perform differential analysis using Cuffdiff and Cuffnorm tool to generate a 17 normalized data set that includes all the samples. FPKM values calculated by Cufflinks were used as 18 input. The transcript quantification engine of Cufflinks, Cuffdiff, was used to calculate transcript 19 expression levels in more than one condition and test them for significant differences. To identify a 20 transcript as being differentially expressed, Cuffdiff tests the observed log-fold-change in its 21 expression against the null hypothesis of no change (i.e. the true log-fold-change is zero). Because of 22 measurement errors, technical variability, and cross-replicate biological variability might result in an 23 observed log-fold-change that is non-zero, Cuffdiff assesses significance using a model of variability 24 in the log-fold-change under the null hypothesis. This model is described in details in (21). The

1 differential gene expression analysis was performed using DESeq (22) and edgeR (23) within the R 2 Bioconductor packages. Genes were considered differentially expressed between two group if they 3 met the following requirement: fold change (FC) > ± 1.5 , p<0.05, FDR <0.05.

Enrichment of gene ontology (GO). GO enrichment analysis amongst differentially expressed genes
(DEGs) was performed using Goseq (24) against the background of full human genome (hg19). GO-

6 terms with adjusted p value < 0.05 were considered significantly enriched.

Clustering of DEGs. We categorized the DEGs according to their response upon silencing of siSTAT2 and siIRF9; categories are listed as A to I (Figure 2E). Then to determine relationship between these categories, we calculated the distance of centers of different categories. For each gene, we transformed siSTAT2 and siIRF9 fold changes (FC) to deviation from the mean FC of the category the respective gene belongs: $FC_{new} = FC_{old} - \varepsilon (FC_{category})$. The parameter ε was estimated to give the perfect match between predefined categories (A to I) and clustering based on Euclidean distance. Results were plotted as a heatmap.

14 Modular transcription analysis. The tmod package in R (25) was used for modular transcription 15 analysis. In brief, each transcriptional module is a set of genes, which shows coherent expression 16 across many biological samples (26, 27). Modular transcription analysis then calculates significant 17 enrichment of a set of foreground genes, here DEGs, in pre-defined transcriptional module compared 18 to a reference set. For transcriptional modules, we used a combined list of 606 distinct functional 19 modules encompassing 12712 genes, defined by Chaussabel et al. (28) and Li et al. (29), as the 20 reference set in tmod package (Supplemental table S5). The hypergeometric test devised in 21 tmodHGtest was used to calculate enrichments and p-values employing Benjamini-Hochberg 22 correction (30) for multiple sampling. All the statistical analyses and graphical presentations were in 23 performed in R (31).

1 Luciferase gene reporter assay

2 U3A or U3A-STAT1 cells at 90% confluency were cotransfected with 100 ng of one of the 3 following CXCL10 promoter containing firefly luciferase reporter plasmids (generously donated by 4 Dr. David Proud, Calgary, (32)), CXCL10prom-972pb-pGL4 (full length -875/+97 promoter), 5 CXCL10prom-376pb-pGL4 (truncated -279/+97 promoter), CXCL10prom972pb-ΔISRE(3)-pGL4 6 (full length promoter with ISRE(3) site mutated), CXCL10prom972pb- $\Delta\kappa B(1)$ -pGL4 (full length 7 promoter with NF κ B(1) site mutated), CXCL10prom972pb- $\Delta\kappa$ B(2)-pGL4 (full length promoter with 8 NF κ B(2) site mutated), CXCL10prom972pb- Δ AP-1-pGL4 (full length promoter with AP-1 site 9 mutated) together with 50ng of pRL-null renilla-luciferase expressing plasmid (internal control). 10 Transfection was performed using Lipofectamine 2000 (Life technologies) using a 1:2 DNA to 11 lipofectamine ratio. At 8 h post-transfection, cells were stimulated for 16 h with either IFNB or 12 IFN β +TNF α . Firefly and renilla luciferase activities were quantified using the Dual-luciferase 13 reporter assay system (Promega). Luciferase activities were calculated as the luciferase/renilla ratio 14 and were expressed as fold over the non-stimulated condition.

15

16 Virus titration by plaque assay

17 Quantification of VSV infectious virions was achieved through methylcellulose plaque forming unit 18 assays. U3A and U3A-STAT1 cells were either left untreated or stimulated with IFNB or 19 IFN β +TNF α for 30h. Cells were then infected with Vesicular Stomatitis Virus (VSV)-GFP (kindly 20 provided by Dr. J. Bell, University of Ottawa, Canada) at an MOI of 5 for 1h in serum free medium 21 (SFM). Cells were then washed twice with SFM and further cultured in DMEM medium containing 22 2% HI-FCl. The supernatants were harvested at 12h post-infection and serial dilutions were used to 23 infect confluent Vero cells (ATCC) for 1h in SFM. The medium was then replaced with 1% 24 methylcellulose in DMEM containing 10% HI-FCl. Two days post-infection, GFP-positive plaques

were detected using a Typhoon Trio apparatus and quantified using the Imagequant software
 (Molecular Dynamics).

3

4 Statistical analyses

5 Statistical analyses of qRT-PCR and luciferase assay results were performed using the Prism 7

- 6 software (GraphPad) using the tests indicated in the figure legends. Statistical significance was
- 7 evaluated using the following P values: P < 0.05 (*), P < 0.01 (**), P < 0.001 (***) or
- 8 P < 0.0001 (****). Differences with a *P*-value < 0.05 were considered significant. Statistical analysis
- 9 of the RNASeq data is described in the Bioinformatics analysis section above.

10

1 **RESULTS**

2

3 Distinct induction profiles of antiviral and immunoregulatory genes in response to IFNβ, TNFα 4 and IFNβ+TNFα.

5 First, we sought to determine the induction profile of a selected panel of immunoregulatory 6 and antiviral genes in response to IFN β +TNF α in comparison to IFN β or TNF α alone. A549 cells 7 were stimulated either with IFN β , TNF α or IFN β +TNF α for various times between 3-24h and the 8 relative mRNA expression levels were quantified by qRT-PCR. Analysis of the expression of the 9 selected genes revealed distinct profiles of response to IFN β , TNF α or IFN β +TNF α (Figure 1). 10 IDO, DUOX2, CXCL10, APOBEC3G, ISG20 and IL33 exhibited synergistic induction in response to 11 IFN β +TNF α compared to IFN β or TNF α alone. Expression in response to IFN β +TNF α increased 12 over time, with maximum expression levels observed between 16 and 24h. While NOD2 and IRF1 13 induction following stimulation with IFN β +TNF α was also significantly higher than upon IFN β or 14 TNF α single cytokine stimulation, they exhibited a steady-state induction profile starting as early as 15 3h. MX1 and PKR, two typical IFN β -inducible ISGs, were found induced by IFN β +TNF α similarly 16 to IFN β alone and were not responsive to TNF α . CCL20 responded to IFN β +TNF α with a kinetic 17 and amplitude similar to TNF α , but was not responsive to IFN β alone. IL8 expression was not 18 induced by IFN β , but was increased by TNF α starting at 3h and remained steady until 24h. In 19 contrast to other genes, *IL8* induction in response to IFN β +TNF α was significantly decreased 20 compared to TNF α alone. Overall, these results confirm that induction of a subset of antiviral and 21 immunoregulatory genes is greatly increased in response to IFN β +TNF α compared to either IFN β or 22 TNF α alone.

23

1 Workflow for genome-wide characterization of the delayed transcriptional program induced

2 by IFN β +TNF α in the absence of STAT1.

In a previous report, we provided evidence of a STAT1-independent, but STAT2- and IRF9-3 4 dependent, pathway engaged downstream of IFN β +TNF α (12). Here, taking advantage of the 5 STAT1-deficient human U3A cell line (13), we aimed to fully characterize the STAT1-independent 6 transcriptional program induced by IFN β +TNF α . Two hallmarks of STAT2 and IRF9 activation, *i.e.* 7 STAT2 Tyr690 phosphorylation and induction of IRF9, were observed in U3A cells following 8 stimulation with IFN β +TNF α . Although STAT2 and IRF9 activation was reduced compared to the 9 parental 2ftGH cells expressing endogenous STAT1, this observation supports the capacity of 10 STAT2 and IRF9 to be activated in STAT1-deficient U3A cells stimulated with IFN β +TNF α 11 (Figure 2A).

12

13 To investigate the transcriptional program triggered independently of STAT1, the U3A cells 14 were efficiently transfected with Control (Ctrl)-, STAT2- or IRF9-RNAi and further left untreated or 15 stimulated with IFN β +TNF α for 16h (Figure 2B). Efficient silencing was confirmed by immunoblot 16 (Figure 2C). To perform genome-wide transcriptional analysis, total RNA was isolated and analyzed 17 by RNA sequencing (n=3 for each group detailed in **Figure 2B**) on an Illumina HiSeq2500 platform. 18 To validate the RNASeq data, the fold changes (FC) of 13 genes between the different experimental 19 groups, siCTRL non-stimulated (NS) vs siCTRL IFN β +TNF α , siCTRL IFN β +TNF α vs siSTAT2 20 IFN β +TNF α and siCTRL IFN β +TNF α vs siIRF9 IFN β +TNF α was confirmed by qRT-PCR. A 21 positive linear relationship between RNASeq and qRT-PCR FC was observed (Figure 2D).

22

An antiviral and immunoregulatory transcriptional signature is induced by IFNβ+TNFα independently of STAT1.

3 To identify STAT1-independent differentially expressed genes (DEGs) upon IFN β +TNF α 4 stimulation, comparison between non-stimulated (NS) and IFN β +TNF α stimulated control cells was 5 performed (Figure 2E). In total, 612 transcripts, including protein-coding transcripts, pseudogenes 6 and long non-coding RNA (lncRNA), were significantly different (FC>1.5, p<0.05, FDR<0.05) in 7 IFN β +TNF α vs NS. Among these, 482 DEGs were upregulated and 130 were downregulated 8 (Figure 3A; See Supplemental Table S1 for a complete list of DEGs). To identify the Biological 9 Processes (BP) and Molecular Functions (MF) induced by IFN β +TNF α independently of STAT1, we 10 first subjected upregulated DEGs through Gene Ontology (GO) enrichment analysis. The top enriched GO BP (p< 10⁻¹⁰) and MF, are depicted in Figure 3B (See Supplemental Table S2 for a 11 12 complete list of enriched GO). The majority of the top enriched BPs were related to cytokine 13 production and function (response to cytokine, cytokine-mediated signaling pathway, cytokine production, regulation of cytokine production), immunoregulation (Immune response, immune 14 15 system process, innate immune response, regulation of immune system process) and host defense 16 response (defense response, response to other organism, 2'-5'-oligoadenylate synthetase activity and 17 dsRNA binding). Fourteen MF categories were found enriched in IFN β +TNF α . The top enriched 18 MFs were related to cytokine and chemokine functions (cytokine activity, cytokine receptor binding, 19 chemokine activity, Interleukin 1-receptor binding). Other enriched MF included peptidase related 20 functions (endopeptidase inhibitor activity, peptidase regulator activity, serine-type endopeptidase 21 activity).

22

23 To further gain insight into the functional significance of the STAT1-independent 24 IFN β +TNF α -induced transcriptional response, we conducted a modular transcription analysis of

1 upregulated DEGs against 606 immune-related functional modules. These modules were previously 2 defined from co-clustered gene sets built via an unbiased data-driven approach as detailed in the 3 material and methods section (28, 29). STAT1-independent IFN β +TNF α -induced DEGs showed 4 significant enrichment in 37 modules (Supplemental Table S3). Six of these modules were 5 associated with virus sensing/Interferon antiviral response, including LI.M75 (antiviral IFN 6 signature), LI.M68 (RIG-I-like receptor signaling), LI.M127 (type I interferon response), LI.M111.0 7 and LI.M111.1 (IRF2 target network) and LI.M150 (innate antiviral response) (Figure 3C). 8 Additionally, 6 modules were associated with immunoregulatory functions, including LI.M29 9 (proinflammatory cytokines and chemokines), LI.M27.0 and LI.M27.1 (chemokine cluster I and II), 10 LI.M38 (chemokines and receptors), LI.M115 (cytokines receptors cluster) and LI.M37.0 (immune 11 activation - generic cluster) (Figure 3C). Module analysis also showed enriched AP-1 transcription 12 factor-related network modules, LI.M20 and LI.M0, and cell cycle and growth arrest LI.M31 13 module. To confirm the capacity of IFN β +TNF α to trigger an antiviral response independently of 14 STAT1, U3A cells were stimulated with IFN β or IFN β +TNF α and further infected with Vesicular 15 stomatitis Virus (VSV). While VSV replicated similarly in untreated U3A cells and in cells treated 16 with IFN β , treatment with IFN β +TNF α significantly restricted VSV replication (Figure 3D). 17 Importantly, IFN β treatment led to a significant antiviral response when STAT1 expression was 18 restored in U3A cells (U3A-STAT1). In this context, the combination of IFN β +TNF α was more 19 effective in restricting VSV than IFN β alone (**Figure 3D**). Collectively, these data demonstrate that 20 the co-stimulation with IFN β and TNF α induces a broad antiviral and immunoregulatory 21 transcriptional program that is independent of STAT1.

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1 Clustering of STAT1-independent IFNβ+TNFα induced DEGs according to their regulation by

2 STAT2 and IRF9.

3 Next, we sought to assess how the STAT1-independent IFN β +TNF α -induced antiviral and 4 immunoregulatory transcriptional response was regulated by STAT2 and IRF9. To do so, we 5 compared transcripts levels in siSTAT2_IFN β +TNF α vs siCTRL IFN β +TNF α and siIRF9 IFN β +TNF α vs siCTRL IFN β +TNF α conditions (Figure 2E and Supplemental Table S1). 6 7 Volcano plots revealed that a fraction of IFN β +TNF α -induced DEGs were significantly (FC>1.5, 8 p<0.05, FDR<0.05) downregulated or upregulated upon silencing of STAT2 (Figure 4A) or IRF9 9 (Figure 4B). Nine distinct theoretical categories of DEGs can be defined based on their potential 10 individual behavior across siSTAT2 and siIRF9 groups (Categories A-I, Figure 2E): a gene can either be downregulated upon STAT2 and IRF9 silencing, indicative of a positive regulation by 11 12 STAT2 and IRF9 (Categorie A); conversely, a gene negatively regulated by STAT2 and IRF9 would 13 exhibit upregulation upon STAT2 and IRF9 silencing (Categorie B); Genes that do not exhibit 14 significant differential expression in siSTAT2 and siIRF9 groups would be classified as STAT2 and 15 IRF9 independent (Categorie C); IRF9-independent genes could exhibit positive (Categorie D) or 16 negative (Categorie E) regulation by STAT2; conversely, STAT2-independent genes might be 17 positively (Categorie F) or negatively (Category G) regulated by IRF9; lastly, STAT2 and IRF9 18 could have opposite effects on a specific gene regulation (Categorie H and I). Based on a priori 19 clustering of RNASeq data (Figure 4C) and analysis of the expression of 18 genes by qRT-PCR 20 (Figure 4D), we found that STAT1-independent IFN β +TNF α -induced DEGs clustered into only 7 of 21 the 9 possible categories. Amongst the 482 DEGs, 163 genes exhibited either inhibition or 22 upregulation following silencing of STAT2 and/or IRF9 (Category B-G). A large majority of 23 upregulated DEGs, i.e. 319 out of the 482 DEGs, were not significantly affected by either STAT2 or 24 IRF9 silencing, and were therefore classified as STAT2/IRF9-independent (Figure 4C). No genes

1 were found in categorie H and only one gene was found in categorie I.

2

3 To functionally interpret these clusters, we applied the modular transcription analysis to each 4 of the categories to assess for the specific enrichment of the functional modules found associated 5 with IFN β +TNF α -upregulated DEGs (Figure 4E). First, most modules, except LI.M31 (cell cycle 6 and growth arrest), LI.M38 (chemokines and receptors), LI.M37.0 (immune activation - generic 7 cluster) and LI.M53 (inflammasome receptors and signaling), were enriched in the category of DEGs 8 positively regulated by STAT2 and IRF9 (Category A). Conversely, the cluster negatively regulated 9 by STAT2 and IRF9 (Category B) exclusively contains enriched LI.M38 (chemokines and receptors), 10 LI.M37.0 (immune activation - generic cluster) and LI.M115 (cytokines receptors cluster). The 11 cluster of IRF9-independent genes that are negatively regulated by STAT2 (Category E) only 12 exhibited enrichment in the virus sensing/ IRF2 target network LI.M111.0 module, while the IRF9-13 independent/ STAT2-positively regulated cluster (Category D) encompasses antiviral and 14 immunoregulatory functions. The STAT2-independent, but IRF9 positively regulated transcripts 15 (Category F) was mainly enriched in modules associated with the IFN antiviral response, including 16 LI.M75 (antiviral IFN signature), LI.M68 (RIG-I-like receptor signaling), LI.M127 (type I interferon 17 response), and LI.M150 (innate antiviral response). Finally, the STAT2-independent, but IRF9 18 negatively regulated cluster (Category G) was mostly enriched in modules associated with 19 immunoregulatory functions, including LI.M29 (proinflammatory cytokines and chemokines), 20 LI.M27.0 and LI.M27.1 (chemokine cluster I and II), LI.M78 (myeloid cell cytokines), but also with 21 cell cycle and growth arrest (LI.M31) and inflammasome receptors and signaling (LI.M53). Of note, 22 all modules were found enriched in the cluster of genes induced independently of STAT2 and IRF9 23 (Category C), pointing to a broad function of this yet to be defined pathway(s) in the regulation of the 24 antiviral and immunoregulatory program elicited by IFN β +TNF α . Altogether these observations

1 reveal that STAT2 and IRF9 are involved in the regulation of only a subset of the genes induced in 2 response to the co-stimulation by IFN β and TNF α in the absence of STAT1. Importantly, our results 3 also reveal that STAT2 and IRF9 act in a concerted manner to regulate a specific subset of the

- 4 IFN β +TNF α -induced DEGs, but are also independently involved in distinct non-canonical pathways.
- 5

6 Differential regulation of *CXCL10* in response to IFN β and IFN β +TNF α .

7 Identification of DEGs upregulated by IFN β +TNF α in a STAT1-independent, but STAT2 and 8 IRF9-dependent, manner potentially reflects the regulation by an alternative STAT2/IRF9-containing 9 complex (4, 7). Whether this STAT2/IRF9 pathway ultimately leads to gene regulation through the 10 same ISRE sites used by the ISGF3 complex remained to be assessed. In our RNASeq analysis 11 (Supplemental Table S1) and qRT-PCR validation (Figure 4D), CXCL10 belongs to the 12 IFN β +TNF α -induced DEGs that are dependent on STAT2 and IRF9. The CXCL10 promoter contains 13 3 ISRE sites. We used full-length wild-type (972bp containing the 3 ISRE sites), truncated (376bp 14 containing only the ISRE(3) site) or mutated (972bp containing a mutated ISRE(3) site) CXCL10 15 promoter luciferase (CXCL10prom-Luc) reporter constructs (Figure 5A) to determine the ISRE 16 consensus site(s) requirement. U3A and STAT1-rescued U3A-STAT1 cells were transfected with the 17 CXCL10 prom-Luc constructs and further stimulated with IFN β or IFN β +TNF α to monitor the 18 canonical ISGF3-dependent and the STAT1-independent responses. In the absence of STAT1, IFNB 19 failed to activate CXCL10prom. However, the induction of the CXCL10prom activity was restored 20 either upon expression of STAT1 or when TNF α was used together with IFN β (Figure 5B). While 21 the activation of CXCL10 prom in response to IFN β in the presence of STAT1 involves the distal 22 ISRE(1) and/or ISRE(2) sites and the proximal ISRE(3) site, only the ISRE(3) site is required for 23 induction by IFN β +TNF α in the absence of STAT1. Additionally, we also assessed the contribution 24 of the two NF- κ B and the AP-1 sites present in the promoter using CXCL10 prom-Luc mutated

1	constructs in comparison to the wild-type reporter. While none of the NF- κ B and the AP-1 sites were
2	required for induction of the CXCL10 promoter by IFN β in the presence of STAT1, the two NF- κ B
3	sites were necessary for the STAT1-independent induction in response to IFN β +TNF α (Figure 5B).
4	These observations suggest that the ISRE site usage is more restricted in the absence of STAT1 in the
5	context of the co-stimulation with IFN β TNF α than in the context of an ISGF3-dependent regulation.
6	
7	DISCUSSION

8

9 Our study was designed to determine the functional relevance of a STAT1-independent, but 10 STAT2- and IRF9-dependent, signaling pathway in the transcriptional program induced by IFN β in 11 the presence of TNF α . Previous studies reported that IFN β and TNF α synergize to induce a specific 12 delayed antiviral program that differs from the response induced by IFN β only. This specific 13 synergy-dependent antiviral response is required for a complete abrogation of Myxoma virus in 14 fibroblasts (10) and contributes to the establishment of a sustained type I and type III IFNs response 15 during paramyxovirus infection in airway epithelial cells (12). The underlying mechanisms of this 16 specific response remain ill defined, but their elucidation is of particular importance with regards to 17 conditions with elevated levels of both IFN β and TNF α such as pathogen intrusion or, autoimmune 18 or chronic inflammatory diseases (8).

19

First, we confirmed the previously documented synergistic induction of an IFN β +TNF α mediated delayed transcriptional program composed of genes that are either not responsive to IFN β or TNF α separately or are only responsive to either one of the cytokine when used separately albeit to a lesser extent (**Figure 1**). Using genome wide RNA sequencing, we demonstrate that 1 IFNβ+TNFα induces a broad transcriptional program in cells deficient in STAT1. GO enrichment 2 and transcriptional module analyses showed that STAT1-independent upregulated DEGs encompass 3 a wide range of immunoregulatory and host defense, mainly antiviral, functions. The STAT1-4 independent antiviral response mounted by STAT1-deficient cells in response to IFNβ+TNFα 5 efficiently restricted VSV replication (**Figure 2**). These findings highlight the functional significance 6 of a STAT1-independent response.

7

8 We also focused on deciphering the role of STAT2 and IRF9 in the STAT1-independent 9 transcriptional program elicited in response to IFN β +TNF α . We previously found that IFN β +TNF α 10 induces the DUOX2 gene via a STAT2- and IRF9-dependent pathway in the absence of STAT1 (12). 11 To what extent this pathway contributes to the STAT1-independent response engaged downstream of 12 IFN β +TNF α remained to be addressed. Based on the possible regulation by STAT2 and/or IRF9, 13 IFNβ+TNFα-induced DEGs could theoretically partitioned into 9 different predicted categories 14 (Figure 2E), but we found that they in fact only significantly segregate into 7 categories. 15 Importantly, the distribution of DEGs amongst 7 different categories reflects distinct contributions of 16 STAT2 or IRF9 and highlights the heterogeneity of the mechanisms of regulation of the 17 IFN β +TNF α -induced genes. Importantly, only one anecdotic gene was found in categories implying 18 inverse regulation by STAT2 and IRF9 (categories H and I) pointing to convergent functions of 19 STAT2 and IRF9 when both are engaged in gene regulation. We can rule out that the distinct 20 regulation mechanisms reflect distinct profiles of induction by IFN β +TNF α . For instance CXCL10, 21 *IL33*, *CCL20* and *ISG20* all exhibit synergistic induction by IFN β +TNF α (Figure 1), but are 22 differentially regulated by STAT2 and/or IRF9; while CXCL10 is dependent on STAT2 and IRF9, 23 IL33 is independent on STAT2 and IRF9, and CCL20 and ISG20 are STAT2-independent but IRF9-24 dependent (Figure 4D).

1

2 Consistent with our previous report (12), we found several genes positively regulated by a 3 non-canonical STAT2- and IRF9-dependent, but STAT1-independent, pathway (Category A). DEGs 4 in this category encompass most of the functions induced in response to IFN β +TNF α , with the 5 notable exception of cell cycle and growth arrest and inflammasome and receptor signaling functions. 6 Genes negatively regulated by STAT2 and IRF9 were also identified (Category B). Accumulating 7 evidence point to the formation of an alternative STAT2/IRF9-containing complex mediating gene 8 expression in the absence of STAT1 (33-37). The specificity of the DNA-binding of a STAT2/IRF9 9 complex compared to the ISGF3 complex remains unclear. It was originally found that STAT2/IRF9 10 exhibit only limited DNA-binding affinity for the typical ISRE sequence in the absence of STAT1 11 (33), but association of STAT2 with the promoter of antiviral genes induced by Dengue virus in 12 STAT1-deficient mice was demonstrated by Chromatin immunoprecipitation (38). Here, the CXCL10 13 gene was further analyzed as a paradigm of STAT2- and IRF9-positively regulated gene. Promoter 14 activity analysis unveiled distinct ISRE site usage between IFN β and IFN β +TNF α stimulation 15 (Figure 5). Interestingly, the ISRE site used in response to IFN β +TNF α lies close to the NF- κ B 16 sites that are also specifically engaged in this response. A possible mechanism for the synergistic 17 action of IFN β +TNF α might be related to the previous description of the cooperativity between 18 ISGF3 and NF- κ B in the context of Listeria infection (39, 40). In this context, ISGF3 and NF- κ B 19 tether a complete functional mediator multi-subunit complex that bridges transcription factors with 20 Pol II and initiation and elongation factors to the promoter of antimicrobial genes (39). Both STAT1 21 and STAT2 functionally and physically interact with the mediator (41, 42). Additionally, IL-6 22 induction by NF- κ B inducers was found synergistically enhanced by IFN β . In this model, 23 unphosphorylated STAT2 was proposed to act as a bridge connecting p65 and IRF9 bound to kB and 24 ISRE consensus sites, respectively (43). Therefore, in the absence of STAT1, the synergistic

1 induction of some IFN β +TNF α -induced DEGs might be explained by the interaction between a 2 STAT2/IRF9-containing complex with NF-κB. However, we cannot exclude that interaction with a 3 vet to be identified alternative transcription factor might tether STAT2 and IRF9 to the DNA in the 4 absence of STAT1. Specific regulation of STAT2 and IRF9 might also contribute to the synergistic 5 response elicited by IFN β +TNF α . In a previous study, we observed IRF9 induction and enhanced/extended STAT2 phosphorylation in response to IFN β +TNF α (12). We hypothesized that 6 7 these events likely contribute to the specific activation of the non-canonical STAT2/IRF9-dependent 8 pathway. Remarkably, a similar prolonged STAT2 phosphorylation was observed upon stimulation 9 of STAT1 KO bone marrow-derived murine macrophages with IFNa. In this context, a STAT2/IRF9 10 complex was shown to induce a delayed set of ISGs (37). Taken together, it is reasonable to 11 speculate that in a physiological context, when TNF α is present with IFN β , a signal is elicited to 12 progressively exclude STAT1 from the STAT2/IRF9 complex and favor the non-canonical 13 STAT2/IRF9 pathway to drive a specific delayed response.

14

15 The observation that some IFN β +TNF α -induced genes were independent of IRF9 but 16 dependent on STAT2, either positively or negatively (Categories D and E), in a STAT1 deficient 17 context might reflect the previous observation that STAT2 forms alternative complexes with other STAT members. STAT2 was shown to associate with STAT3 and STAT6, although it is not 18 19 completely clear whether IRF9 is also required in these alternative complexes (44, 45). Interestingly, 20 transcriptional module analyses demonstrated that the functional distribution of genes negatively 21 regulated by STAT2 is very limited compared to other categories; only a virus-sensing module was 22 enriched in this category. In contrary, IRF9-independent genes positively regulated by STAT2 23 mediate broader antiviral and immunoregulatory functions.

1 ISGF3-independent functions of IRF9 have been proposed based on the study of IRF9 2 deficiencies (reviewed in (7, 46)). However, IRF9 target genes in these contexts have been barely 3 documented. Intriguingly, Li et al. (47) studied IFN α -induced genes and their dependency on the 4 ISGF3 subunits. While they confirmed previous studies showing that IFN α can trigger a delayed and 5 sustained ISGs response via an ISGF3-independent pathway, it is very striking that they did not find 6 STAT1- and STAT2-independent but IRF9-dependent genes. Indeed, all identified IRF9-dependent 7 genes were either STAT2- or STAT1-dependent. This result greatly differs with our study. Here, we 8 found several IFN β +TNF α -induced DEGs independent of STAT1 and STAT2, but positively or 9 negatively regulated by IRF9 (Categories F and G). Typically, IRF9 is considered a positive regulator 10 of gene transcription. However, our findings are consistent with recent reports documenting the role 11 of IRF9 in the negative regulation of the TRIF/NF-KB transcriptional response (48) or the expression 12 of SIRT1 in acute myeloid leukemia cells (49). The molecular mechanism underlying gene regulation 13 by IRF9 without association with either STAT1 or STAT2 remain to be elucidated. To the best of our 14 knowledge, no alternative IRF9-containing complex has been described.

15

16 Unexpectedly, a vast majority of genes were found to be independent of STAT2 and IRF9 17 (Categorie C). All transcriptional modules were enriched in this category pointing to a major role of a 18 yet to be identified pathway in the establishment of a host defense and immunoregulatory response. 19 The NF- κ B pathway is widely known to be engaged downstream of the TNF receptor. While NF- κ B 20 is an obvious candidate for being involved in the regulation of the STAT2 and IRF9-independent 21 DEGs, this might fall short in explaining the synergistic action of IFN β +TNF α . Synergistic 22 activation of NF- κ B was reported in the context of IFN γ and TNF α treatment (50). However, we did 23 not observe enhanced NF- κ B activation upon IFN β +TNF α stimulation compared to TNF α alone 24 (12). Alternative pathways might be of interest. For instance, the IL33 gene is synergistically induced

1	by IFN β +TNF α in a STAT2 and IRF9-independent manner; scanning of the <i>IL33</i> promoter for
2	transcription factor binding sites revealed AP-1, NF-kB and IRF7 consensus sites (51). The potential
3	role of AP-1 is also supported by the finding that the AP-1 transcription network module is enriched
4	amongst IFN β +TNF α -induced DEGs (Figure 4E). However, this module is not restricted to genes
5	regulated independently of STAT2 and IRF9. It is also worth noting that two modules of IRF2-target
6	genes were enriched, although again not specifically in the STAT2- and IRF-9 independent category
7	(Figure 4E). Finally, increased colocalized recruitment of IRF1 and p65 to the promoter of a subset
8	of genes induced by IFN α and TNF α in macrophages was observed (52). However, while IRF1 was
9	found synergistically induced by IFN β +TNF α at early stages (Figure 1), we did not observe
10	significant induction of IRF1 in the absence of STAT1 by RNASeq (Supplemental Table S1) or
11	qRT-PCR (data not shown). Further studies will be needed to challenge the role of these various
12	pathways in the synergistic induction of genes by IFN β +TNF α independently of STAT2 and IRF9.

13

Altogether our results demonstrate that in conditions with elevated levels of IFNβ and TNFα, a broad antiviral and immunoregulatory delayed transcriptional program is elicited independently of STAT1. Our findings highlight the importance of diverse non-canonical STAT2 and/or IRF9 pathways. Consistent with the growing literature, IFNβ and TNFα synergistic action is in part mediated by the concerted action of STAT2 and IRF9, most likely present in an alternative complex. Finally, our study reveals specific independent roles of STAT2 and IRF9 in the regulation of distinct sets of IFNβ and TNFα-induced genes.

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11

12 Author Contributions

MM and NG conceived and designed the experiments. MKM, AF, EC, MK, ANH, DK, SCO, and
EM performed experiments. PD and AS performed bioinformatics analysis. MKM and NG analyzed
the data. NG wrote the manuscript. All co-authors edited and approved the manuscript.

16

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23		

1 Figure legends

2	
3	Figure. 1: Expression of a panel of antiviral and immunoregulatory genes in response to IFN β ,
4	TNF α and IFN β +TNF α .
5	A549 cells were stimulated with either TNF α , IFN β or costimulated with IFN β +TNF α for the
6	indicated times. Quantification of mRNA was performed by qRT-PCR and expressed as fold
7	expression after normalization to the S9 mRNA levels using the $\Delta\Delta$ Ct method. Mean +/- SEM, n \geq 3.
8	Statistical comparison of TNF α vs. IFN β +TNF α and IFN β vs. IFN β +TNF α was conducted using
9	two-way ANOVA with Tukey's post-test. $P < 0.01$ (**), $P < 0.001$ (***) or $P < 0.0001$ (****).
10	
11	Figure 2: Experimental design used to study the STAT1-independent delayed transcriptional
12	program induced by the combination of IFN β and TNF α .
13	A) U3A (STAT1-deficient) and 2ftGH (parental STAT1-positive) cells were left untreated or
14	stimulated with IFN β +TNF α for the indicated times. WCE were analyzed by SDS-PAGE followed
15	by immunoblot using anti STAT1-P-Tyr701, total STAT1, STAT2-P-Tyr690, total STAT2, IRF9 or
16	actin antibodies. B-E) U3A cells were transfected with siCTRL, siSTAT2 or siIRF9 before being left
17	untreated (NS) or stimulated with IFN β +TNF α for 24h. In B, The schematic describes the workflow
18	of sample preparation and analysis. In C, WCE were analyzed by SDS-PAGE followed by
19	immunoblot using anti STAT2, IRF9 and actin antibodies. In D, Graph showing the correlation
20	between fold-changes (FC) measured by RNASeq and qRT-PCR for 13 genes. Data from siCTRL
21	NS vs siCTRL IFN β +TNF α , siSTAT2 IFN β +TNF α vs siCTRL IFN β +TNF α , siIRF9 IFN β +TNF α
22	vs siCTRL IFN β +TNF α conditions were used. In E, Diagram describing the bioinformatics analysis
23	strategy used to determine STAT1-independent differentially expressed genes (DEGs) and their
24	regulation by STAT2 and IRF9.

1

2 Figure 3: Functional characterization of STAT1-independent IFNβ+TNFα-induced DEGs.

A) Volcano plots of the fold-change (FC) vs. adjusted P-value of IFNB+TNFa (I+T) vs non-3 4 stimulated (NS) siCtrl conditions. B) Gene ontology (GO) enrichment analysis of the differentially 5 upregulated genes in IFN β +TNF α vs non-stimulated siCtrl conditions based on the Biological Processes and Molecular Function categories. C) Modular transcription analysis of upregulated 6 DEGs. Eighteen enriched modules are shown. The full list of enriched modules is available 7 8 **Supplemental Table S3. D**) U3A and STAT1-rescued U3A-STAT1 cells were stimulated with IFNβ 9 (I) or IFN β +TNF α for 30h before infection with VSV at a MOI of 5 for 12h. The release of 10 infectious viral particles was quantified by plaque forming unit (pfu) assay. The left graphs show dot-11 plots of all stimulations. Statistical comparisons were performed on the "before and after" plots 12 (displayed on the right of dot-plot graphs) using ratio paired t-tests. Top panel shows the immunoblot 13 in 2ftGH, U3A-STAT1 and U3A cells using anti-STAT1 and anti-tubulin (loading control) 14 antibodies.

15

16 Figure 4: Clustering of IFNβ+TNFα-induced DEGs according to their regulation by STAT2 17 and IRF9.

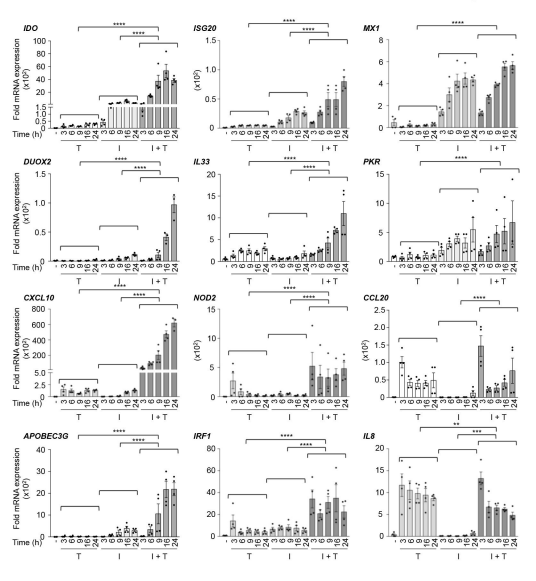
A) Volcano plots of the fold-change *vs.* adjusted *p*-value of siSTAT2 IFN β +TNF α *vs.* siCTRL IFN β +TNF α (I+T) conditions. **B**) Volcano plots of the fold-change *vs.* adjusted *p*-value of siIRF9 IFN β +TNF α *vs.* siCTRL IFN β +TNF α conditions. **C**) Hierarchical clustering of the categories of DEG responses according to their regulation by STAT2 and IRF9. Euclidean distance metric is used for the construction of distance matrix and the categories are used *a priori* input into clustering algorithm as detailed in Materials and Methods. **D**) Validation of DEGs regulation profile by qRT-PCR. U3A cells were transfected with siCTRL (C), siSTAT2 (S2) or siIRF9 (F9). Cells were further

1 left untreated or stimulated with IFNβ+TNFα for 24h. Quantification of the mRNA corresponding to 2 the indicated genes was performed by qRT-PCR and expressed as relative expression (ΔΔCt) after 3 normalization to the S9 mRNA levels. Mean +/- SEM, n≥5. Statistical comparison of siSTAT2 4 IFNβ+TNFα vs siCTRL IFNβ+TNFα and siIRF9 IFNβ+TNFα vs siCTRL IFNβ+TNFα conditions 5 was conducted using one-way ANOVA with Dunett post-test. P < 0.05 (*), P < 0.01 (**), P < 0.0016 (***) or P < 0.0001 (****). E) Diagram showing enriched transcription modules in each gene 7 category.

8

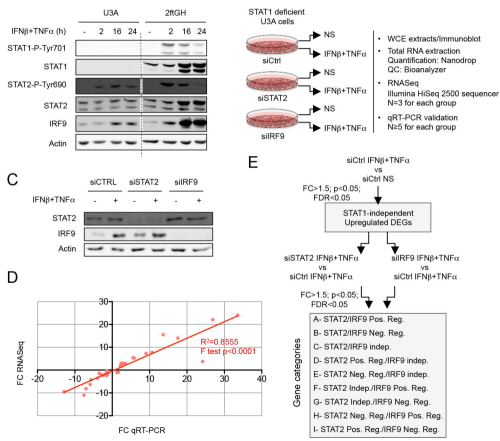
9 Figure 5: Analysis of the *CXCL10* promoter regulation in response to IFN β +TNF α vs IFN β .

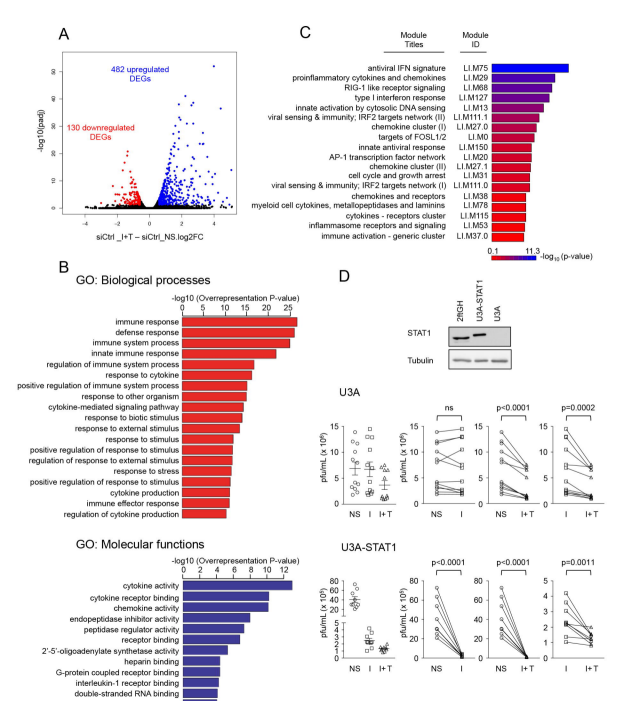
A) Schematic representation of the *CXCL10* promoter (CXCL10prom) luciferase constructs used in this study indicating the main transcription factors consensus binding sites. **B**) U3A and U3A-STAT1 cells were transfected with the indicated CXCL10prom-luciferase reporter constructs and either left untreated or stimulated with IFN β or IFN β +TNF α . Relative luciferase activities were measured at 16h post-stimulation and expressed as fold over the corresponding unstimulated condition. Mean +/-SEM, n=6. Statistical analyses were performed using an unpaired t-test comparing each promoter to the CXCL10prom-972bp construct. *P* < 0.01 (**), *P* < 0.001 (***).



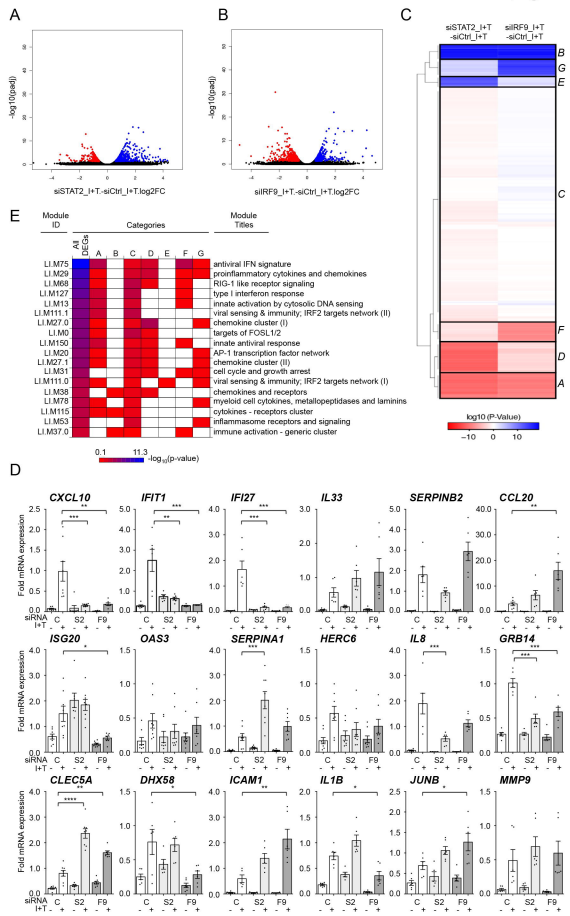
В

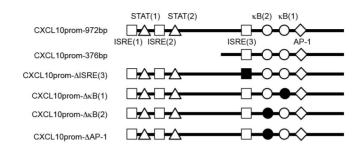
А



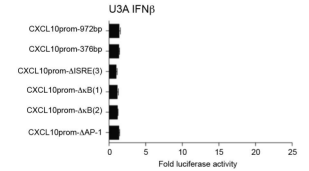


glycosaminoglycan binding collagen binding serine-type endopeptidase activity



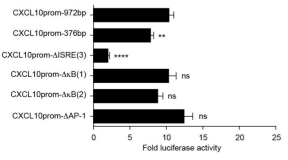




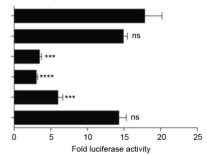


А





U3A IFN β + TNF α



U3A-STAT1 IFN β + TNF α

